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Dec 17 The CA Lexicon available in the CAPLUS and CA files
Feb 06 Engineering Information Encompass files have new names
Feb 16 TOXLINE no longer being updated
Apr 23 Search Derwent WPINDEX by chemical structure
Apr 23 PRE-1967 REFERENCES NOW SEARCHABLE IN CAPLUS AND CA NEWS 3 Feb 06 4 Feb 16 NEWS 5 Apr 23 6 Apr 23 7 May 07 8 Jun 20 NEWS NEWS NEWS DGENE Reload Published patent applications (Al) are now in USPATFULL New SDI alert frequency now available in Derwent's DWPI and DPCI 9 NEWS JUL 13 In-process records and more frequent updates now in NEWS 10 Aug 23 MEDLINE MEDITION PAGE IMAGES FOR 1947-1966 RECORDS IN CAPLUS AND CA Adis Newsletters (ADISNEWS) now available on STN IMSworld Pharmaceutical Company Directory name change to PHARMASEARCH NEWS 11 Aug 23 NEWS 12 Aug 23 NEWS 13 Sep 17 NEWS EXPRESS AUGUST 15 CURRENT WINDOWS VERSION IS V6.0c,
CURRENT MACINTOSH VERSION IS V6.0 (ENG) AND V6.0J (JP),
AND CURRENT DISCOVER FILE IS DATED 07 AUGUST 2001
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COST IN U.S. DOLLARS SINCE FILE TOTAL SESSION ENTRY FULL ESTIMATED COST 0.15 0.15

FILE 'MEDLINE' ENTERED AT 16:20:32 ON 21 SEP 2001

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FILE 'BIOSIS' ENTERED AT 16:20:32 ON 21 SEP 2001 COPYRIGHT (C) 2001 BIOSIS(R)

L1

s haparanase (5N) antibod? 0 HAPARANASE (5N) ANTIBOD?

=> s heparanase (5N) antibod? 1.2

53 HEPARANASE (5N) ANTIBOD?

=> s 12 (10N) (endogenous or natural) L3 0 L2 (10N) (ENDOGENOUS OR NATURAL)

=> dup rem 12 PROCESSING COMPLETED FOR L2

23 DUP REM L2 (30 DUPLICATES REMOVED)

=> dis 14 1-23 ibib abs kwic

ANSWER 1 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

2001:229058 CAPLUS 134:262849

TITLE:

Human heparanase-2, its sequence, recombinant production, and use in identifying potential antagonists and/or agonists Duecker, Klaus; Sirrenberg, Christian Merck Patent G.m.b.H., Germany

INVENTOR(S):

PATENT ASSIGNEE (S): PCT Int. Appl., 46 pp. CODEN: PIXXD2 Patent SOURCE:

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE PATENT NO. APPLICATION NO. DATE

W0 2001021814 A1 20010329 W0 2000-EP8837 20000911 W: CA, JP, US RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

PRIORITY APPLN. INFO.:

EP 1999-118805 A 19990923 EP 2000-114649 A 20000707 20000707

The invention provides a cDNA mol. encoding a human protein believed to be heparanase-2, based on sequence homol. to known heparanases. The

invention also provides polynucleotides that the intain fragments of said cDNA mols. that can be used as hybridization probes or as primers for nucleic acid amplification. The invention further provides expression vectors comprising said cDNA mols., host cells transformed with said vectors for the recombinant prodn. of human heparanase-2. Still further, the invention provides for the use of said human heparanase-2 polypeptides in identifying compds. that may be antagonists and/or agonists, which may be potentially useful in therapy. Finally, the invention provides a fusion protein consisting of said heparanase-2 fused to a Ig Fc region, and antibodies specific for heparanase-2. The cDNA sequence, as well as the corresponding amino acid sequence of human heparanase-2 are claimed. The invention used reverse transcription-polymerase chain reaction (RT-PCR) to show the expression of heparanase-2 gene in various tissues and tumors, and showed the expression of heparanase-2 in transformed 293 human kidney fibroblasts. heparanase-2 in transformed 293 human kidney fibroblasts.
REFERENCE COUNT: 6 (1) Fairbanks, M: WO 9943830 A 1999 CAPLUS REFERENCE (S): (2) Hadasit Med Res Service; WO 9911798 A 1999 CAPLUS (3) Hamdorf, B; WO 9921975 A 1999 CAPLUS Kosir, M; JOURNAL OF SURGICAL RESEARCH 1997, V67(1), P98 CAPLUS (4) KOSIT, M. JOURNAL OF SURGICAL RESEARCH 1997, V67(11), P98 CAPLUS

(5) KOSIT, M. JOURNAL OF SURGICAL RESEARCH 1999, V01(1), P42 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

The invention provides a CDNA mol. encoding a human protein believed to be heparanase-2, based on sequence homol. to known heparanases. The invention also provides polynucleotides that contain fragments of said cDNA mols. that can be used as hybridization probes or as primers for nucleic acid amplification. The invention further provides expression vectors comprising said cDNA mols., host cells transformed with said vectors for the recombinant prodn. of human heparanase-2. Still further, the invention provides for the use of said human heparanase-2 polypeptides in identifying compds. that may be antagonists and/or agonists, which may be potentially useful in therapy. Finally, the invention provides a fusion protein consisting of said heparanase-2 fused to a Ig Fc region, and antibodies specific for heparanase-2. The cDNA sequence, as well as the corresponding amino acid sequence of human heparanase-2 are claimed. The invention used reverse transcription-polymerase chain reaction (RT-PCR) to show the expression of heparanase-2 gene in various tissues and tumors, and showed the expression of heparanase-2. But DIM (Richardical study): MSFS RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (anti-human heparanase 2 specific antibodies) ANSWER 2 OF 23 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 2001:12473 CAPLUS DOCUMENT NUMBER: TITLE: Protein and cDNA sequences of a novel human heparanase gene hnhpl and its splicing variants
Pecker, Iris; Michal, Israel; Itzhaki, Hanan
Insight Strategy & Marketing Ltd., Israel
PCT Int. Appl., 67 pp.
CODEN: PIXXD2 INVENTOR (S): PATENT ASSIGNEE (S): DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. WO 2001000643 A2 20010104 WO 2000-IL358 20000619 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HH, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM SU, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO:

B The invention provides protein and cDNA sequences of a novel human heparanase gene hnhpl and two variants resulted from alternative splicing. The longest clone is 2060 nucleotide long and it contains an open reading frame of 1776 nucleotides, which encodes a polypeptide of 592 amino acids, with a calcd. mol. wt. of 66.5 kDa. The two shorter forms contain an in frame deletion as a result of alternative splicing, one is 162 nucleotides (nt473-630) corresponding to amino acids 150-203, and one is 336 nucleotides (nt473-808) corresponding to amino acids 150-261. The hnhpl gene is mapped to chromosome 10, next to the marker SHGC-57721. The tissue distribution of hnhpl transcripts is detd. The invention also relates to constructing hnhpl gene expression vector to produce recombinant proteins in mammalian cells, which may have heparanase or other glycosyl hydrolase activity, its antibodies, and antisense oligonucleotide and ribozymes for gene modulation and therapeutic use.

REFERENCE (S): (2) Bashkin, P; Biochemistry 1989, V28, P1737 CAPLUS (2) Bashkin, P; Biochemistry 1989, V28, P1737 CAPLUS(3) Burgess, W; Annu Rev Biochem 1989, V58, P575 REFERENCE (S): CAPLUS (5) Chen, Y; Nature Medicine 1997, V3, P866 CAPLUS (7) Durand, P; Glycobiology 1997, V7(2), P277 CAPLUS (8) Eisenberg, S; J Clin Invest 1992, V90, P2013 ALL CITATIONS AVAILABLE IN THE RE FORMAT ARL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
(to heparanase encoded by gene hnhpl; protein and cDNA sequences of a novel human heparanase gene hnhpl and splicing variants) ANSWER 3 OF 23 CAPLUS COPYRIGHT 2001 ACS CAPLUS 2001:57239 134:128217

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: 1.34:1.28217
Heparanase specific molecular probes and their use in research and medical applications
Pecker, Iris; Vlodavsky, Israel; Friedman, Yael;
Perets, Tuvia
Insight Strategy & Marketing Ltd., Israel
U.S., 41 pp., Cont.-in-part of U.S. 5,960,822.
CODEN: USXXAM
Patent INVENTOR (S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE:

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LANGUAGE:
                                                                                   English
 FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                 PATENT NO.
                                                                        KIND DATE
                                                                                                                                             APPLICATION NO. DATE
                                                                                                                                             US 1998-71739
US 1997-922170
                 US 6177545
                                                                                                                                                                                                      19980501
                 US 5968822
                                                                                            19991019
                                                                                                                                                                                                      19970902
               W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, UI, ID, IL, IN, IS, MN, MM, MX, NO, NZ, PL, FT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, FT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 9938706. Al 1999123

AU 1999-38706 19990429

RI AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, NL, SE, IE, FI NO 990629 A 20000224

RITY APPLN. INFO: US 1997-922170 A2 19970902

US 1997-922170 A2 19980511
                                                                                           19991111
                                                                                                                                              WO 1999-US9255
                                                                                                                                                                                                     19990429
                                                                                                                                                                                       A2 19970902
A 19980501
W 19990429
 PRIORITY APPLN. INFO.:
             US 1998-71739 A 19980501
W0 1999-US9255 W 19990429
A variety of heparanase specific mol. probes which can be used for research and medical applications including diagnosis and therapy. Specific applications include the use of a heparanase specific mol. probe for detection of the presence, absence or level of heparanase expression; the use of a heparanase specific mol. probe for therapy of a condition assocd. With expression of heparanase; the use of a heparanase pocific mol. probe for quantification of heparanase in a body fluid: the use of a
mol. probe for quantification of heparanase in a body fluid; the use of a heparanase specific mol. probe for targeted drug delivery; and the use of a heparanase specific mol. probe as a therapeutic agent.

REFERENCE COUNT:

23

REFERENCE(S):

(1) Burgess; Annu Rev Blochem 1989, V58, P575 CAPLUS
                                                                                   (1) Burgess; Annu Rev Biochem 1989, V58, P575 CAPLUS
(4) Folkman; Science 1987, V235, P442 CAPLUS
(6) Fuks; US 5362641 1994 CAPLUS
                                                                                    (9)
                                                                                              Jackson; Physiological Reviews 1991, V71(2), P481
                (10) Kjellen; Annu Rev Biochem 1991, V60, P443 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
heparanase probe antibody; sequence gene human
ST
                  heparanase
                 Antibodies
RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
(heparanase-specific mol. probes and their use in research
                           and medical applications)
                 Antibodies
                 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (monoclonal; heparanase-specific mol. probes and their use in research and medical applications)
 L4 ANSWER 4 OF 23
ACCESSION NUMBER:
                                                                 MEDLINE
2001252356
                                                                                                                                                                                         DUPLICATE 1
                                                                                                                 MEDLINE
                                                                 2001252356 MEDLINE
21248515 PubMed ID: 11350898
The clinicopathological significance of heparanase and basic fibroblast growth factor expressions in hepatocellular carcinoma.
El-Assal O N; Yamanoi A; Ono T; Kohno H; Nagasue N
The Second Department of Surgery, Shimane Medical University, Izumo 693-8501, Japan.
CLINICAL CANCER RESEARCH, (2001 May) 7 (5) 1299-305.
 DOCUMENT NUMBER:
 TITLE:
 AUTHOR:
 CORPORATE SOURCE:
 SOURCE:
                                                                    Journal code: C2H; 9502500. ISSN: 1078-0432
 PUB. COUNTRY:
                                                                   United States
                                                                   Journal; Article; (JOURNAL ARTICLE)
 LANGUAGE:
                                                                   English
 FILE SEGMENT:
ENTRY MONTH:
                                                                   Priority Journals
200107
                                                                   Entered STN: 20010723
               Y DATE:

Entered STN: 20010723

Last Updated on STN: 20010723

Entered Medline: 20010719

Heparan sulfate plays an essential role for insolubility of the components of extracellular matrix and represents a storage depot for various growth factors. Therefore, heparanase produced by a given tumor may facilitate tumor invasiveness and angiogenesis through the release of heparan sulfate-bound growth factors. Although the growth factors responsible for angiogenesis in hepatocellular carcinoma (HCC) have recently been investigated, the clinicopathological significance of heparanase in connection with basic fibroblast growth factor (bFGF) expression in HCC has not been evaluated so far. Fifty-five patients who had undergone hepatic resection for HCC without preoperative treatment were included in the present study. Expression of heparanase mRNA was evaluated by reverse
 ENTRY DATE:
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hepatic resection for HCC without preoperative treatment were included in the present study. Expression of heparanase mRNA was evaluated by reverse transcription-PCR, and bFGF was examined by Western blotting using a monoclonal antibody. Tumor angiogenesis was evaluated by immunostaining with a factor VIII-related monoclonal antibody. Expression of heparanase mRNA was detected in 47% of HCCs and was significantly correlated with larger tumor size (P = 0.01), presence of portal vein invasion (P = 0.01), and higher overall tumor invasiveness (P = 0.02). Moreover, its expression was correlated with tumor microvessel density (MVD; P = 0.02). There was a direct correlation between the levels of bFGF proteins and MVD in HCCs (P = 0.001), and, furthermore, concomitant expression of bFGF and heparanase was associated with higher tumor MVD as compared with expression of either factor alone (P = 0.01). In conclusion, the expression of heparanase in HCC enhances growth, invasion, and angiogenesis of the tumor, and bFGF seems to be a potent angiogenic factor for HCC.

was examined by Western blotting using a monoclonal antibody. Tumor angiogenesis was evaluated by immunostaining with a factor VIII-related monoclonal antibody. Expression of heparanase mRNA was detected in 47% of HCCs and was significantly correlated with larger tumor size (P = 0.01), presence of. .

DUPLICATE 2 ANSWER 5 OF 23 MEDLINE ACCESSION NUMBER: DOCUMENT NUMBER: 2001252533 MEDLINE 21248752 PubMed ID: 11351242 TITLE: Expression of heparanase, Mdm2, and erbB2 in ovarian

AUTHOR: Ginath S: Menczer J: Friedmann Y: Aingorn H: Aviv A: Tajima

for HCC.

K; Dantes A; Glezerman M; Vlodavsky I; Amsterdam A Department of Molecular Cell Biology, Weizmann Institute of CORPORATE SOURCE: Science, Rehovot,

INTERNATIONAL JOURNAL OF ONCOLOGY, (2001 Jun) 18 (6) SOURCE:

1133-44.

Journal code: CX5; 9306042. ISSN: 1019-6439.

PUB. COUNTRY: Greece

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: ENTRY MONTH: Priority Journals 200107

Entered STN: 20010730 ENTRY DATE:

Last Updated on STN: 20010730 Entered Medline: 20010726

Last Updated on STN: 20010730
Entered Medline: 20010726

Ovarian cancer is the most lethal of gynecological malignancies. Yet early diagnosis and prognosis are far from being satisfactory. Degradation of heparan sulfate proteoglycans by heparanase appears to play an important role in the invasiveness of tumor cells through the basement membrane and into the extracellular matrix. Recent cloning of the haparanase gene and generation of monoclonal antibodies against the enzyme permit to examine tumor cell expression of the enzyme. The aim of the present study was to assess heparanase activity and localization in various subtypes of epithelial ovarian cancer in correlation with oncogene expression. Histologically confirmed malignant ovarian tissue from ten women and tissue from 2 benign ovarian tumors and 4 normal ovaries were assessed for heparanase presence, activity and localization, incidence of apoptosis and expression of the oncogenes erbB2 and Mdm2. Heparanase immunohistostaining and activity were present in mucinous carcinomas and were more intense than in endometrioid and in serous carcinomas. The lowest activity was observed in benign ovarian tumors and normal ovaries. In ovarian carcinomas the enzyme was intensely concentrated in the cytoplasm of the cancerous cells. In contrast, in normal ovaries and benign tumors the enzyme was predominantly localized in endothelial cells lining blood capillaries. The rate of apoptosis was considerably higher in mucinous and endometrioid carcinomas, and was lower in serous and primary peritoneal carcinomas. Extremely high concentration of heparanase was often demonstrated in apoptotic cells. Endometrioid and serous carcinomas showed low expression. In benign ovarian tumors and normal ovaries the expression of both oncoproteins was extremely low. In conclusion ovarian carcinomas demonstrate higher levels of heparanase than benign tumors and normal ovaries suggesting that the enzyme may play an important role in tumor progression, expression of oncogenes, such as erbB2 a

ANSWER 6 OF 23 CAPLUS COPYRIGHT 2001 ACS SSION NUMBER: 2000:900837 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

134:38857 TITLE:

Heparanase assay using an immobilized heparan sulfate glycosaminoglycan and a paracrine cell regulator Brenchley, Paul Ernest Charles

INVENTOR (S): Central Manchester Healthcare NHS Trust, UK

PATENT ASSIGNEE(S): PCT Int. Appl., 21 pp. CODEN: PIXXD2

Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 2000077241 A2 20001221 WO 2000-GB2117 20000612 2000077241 20010322 A3

AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO:

BB 1999-13415 A 19996610

ARITY APPLN. INFO.:

GB 1999-13415 A 19990610

A method of assaying a sample to det. heparanase activity thereof comprises the steps of (i) incubating the sample in the presence of a first solid phase support having immobilized thereon an heparan sulfate glycosaminoglycan (HSGAG) polymer substrate for the heparanase, the said substrate being insensitive to the action of proteases and the said substrate having bonded thereto a first binding moiety and having further bonded thereto a paracrine cell regulator (such as cytokine, chemokine or growth factor) capable of binding to HSGAG, (ii) treating the incubated sample with a second solid phase support having a second moiety provided thereon capable of immobilizing HSGAG polymer substrate cleaved from the first solid phase support on said second solid phase support by binding of said second moiety either with the paracrine cell regulator or with the first or second moiety of the cleaved substrate immobilized in the second support solid phase, and (iv) measuring the signal on the second solid phase support.

solid phase support sepd. from the first solid phase support. 9003-99-0, Peroxidase RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (antibody coupled with; heparanase detn. using immobilized heparan sulfate glycosaminoglycan and paracrine cell regulator

ANSWER 7 OF 23 CAPLUS COPYRIGHT 2001 ACS SSION NUMBER: 2000:314574 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 132:333392

Heparanase activity neutralizing anti-heparanase monoclonal antibody Peretz, Tuvia; Miron, Daphna; Shlomi, Yinon; Pecker, Iris; Ayal-Hershkovitz, Maty; Friedman, Yael; INVENTOR(S):

Vlodavsky, Israel

PATENT ASSIGNEE (S): Insight Strategy & Marketing Ltd., Israel; Hadasit Medical Research Services & Development Ltd.;

Friedman, Mark M. PCT Int. Appl., 28 pp. CODEN: PIXXD2

SOURCE:

DOCUMENT TYPE:

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English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
      PATENT NO.
                           KIND DATE
      WO 2000025817
                                  20000511
                            A1
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WC 2000025817 A1 20000511 WC 1999-US25451 19991028

W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MM, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1126878 A1 20010829 EP 1999-956781 19991028

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
NO 2001002190 A 20010612 NO 2001-2190 20010503

RITY APPLN. INFO:

NO 2001-2190 20010503 US 1998-186200 A 19981104 WO 1999-US25451 W 19991028 PRIORITY APPLN. INFO.:

WO 1999-US25451 W 19991028
A monoclonal antibody elicited by a heparanase protein
or an immunogenic portion thereof, the monoclonal antibody
specifically inhibits heparanase activity. The
heparanase-specific monoclonal antibody may be human or
humanized antibody and is useful for treating conditions assocd. with
altered function of a heparan sulfate proteoglycan-assocd biol. effector
mol. such as growth factor, chemokine, cytokine and degradative enzyme.
The condition is selected from the group consisting of angiogenesis, cell
proliferation, tumor, metastasis, inflammatory disorders and autoimmune
conditions.

conditions. REFERENCE COUNT: REFERENCE(S):

(1) Anon; WO 9201003 A1 1992 CAPLUS (2) Fuks; US 5362641 A 1994 CAPLUS (3) Jin, L; Internat J Cancer 1990, V45, P1088 CAPLUS (4) Lider, O; Eur J Immunol 1990, V20, P493 CAPLUS (5) Parish, C; Immunol Cell Biol 1998, V76, P104

APPLICATION NO. DATE

19991028

WO 1999-US25451

CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

Heparanase activity neutralizing anti-heparanase TI

Heparanase activity neutralizing anti-heparanase monoclonal antibody
A monoclonal antibody elicited by a heparanase protein or an immunogenic portion thereof, the monoclonal antibody specifically inhibits heparanase activity. The heparanase-specific monoclonal antibody may be human or humanized antibody and is useful for treating conditions assocd. With altered function of a heparan sulfate proteoglycan-assocd biol. effector mol. such as growth factor, chemokine, cytokine and degradative enzyme. The condition is selected from the group consisting of angiogenesis, cell proliferation, tumor, metastasis, inflammatory disorders and autoimmune conditions.
heparanase neutralizing monoclonal antibody tumor AB

heparanase neutralizing monoclonal antibody tumor inflammation; autoimmune disease angiogenesis heparanase ST

monoclonal antibody ΙT

monoclonal antibody
Chemokines
RL: ADV (Adverse effect, including toxicity); BSU (Biological study,
unclassified); BIOL (Biological study)
(IP-10; anti-heparanase monoclonal antibody for
treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)

Chemokines

KL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study) (MGSA; anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)

Chemokines

Chemoxines
RL: ADV (Adverse effect, including toxicity); BSU (Biological study,
unclassified); BIOL (Biological study)
(PF-4; anti-heparanase monoclonal antibody for
treating angiogenic conditions, tumors, inflammations, metastasis and

autoimmune diseases) Autoimmune disease

Drugs Hybridoma Inflammation

(anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)

Chemokines

Cytokines Growth factors, animal

Interleukin 3 Interleukin 8

Lymphotoxin
Macrophage inflammatory protein 1.alpha.
Macrophage inflammatory protein 1.beta.
Macrophage inflammatory protein 1.beta.
Monocyte chemoattractant protein-1
Neutrophil-activating peptide-2
RANTES (chemokine)
Tumor necrosis factors
RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study)
(anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)
Enzymes, biological studies

Enzymes, biological studies
RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study)
(degradative; anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)

Angiogenesis
(disease; anti-heparanase monoclonal antibody for
treating angiogenic conditions, tumors, inflammations, metastasis and
autoimmune diseases)
Proteoglycans, biological studies

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (heparitin sulfate-contg.; anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors,

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inflammations, metastasis and autoimmu
IT
            Epitopes
                   (mapping; anti-heparanase monoclonal antibody for . treating angiogenic conditions, tumors, inflammations, metastasis and
                   autoimmune diseases)
                    (metastasis; anti-heparanase monoclonal antibody
                   for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune diseases)
            Antibodies
ΙT
            Antibodies
RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
(monoclonal, human or humanized; anti-heparanase monoclonal
antibody for treating anglogenic conditions, tumors,
                    inflammations, metastasis and autoimmune diseases)
            Antibodies
           Antibodies
RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
(monoclonal, neutralizing; anti-heparanase monoclonal
antibody for treating angiogenic conditions, tumors,
inflammations, metastasis and autoimmune diseases)
           Disease, animal (proliferative; anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis
                    and autoimmune diseases)
ΙT
            Interferons
            RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study)
                    (.gamma.; anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and
                    autoimmune diseases)
           autorimiume diseases)
89800-66-8, Heparanase
RL: BSU (Biological study, unclassified); BIOL (Biological study)
(anti-heparanase monoclonal antibody for treating
angiogenic conditions, tumors, inflammations, metastasis and autoimmune
           Giseases) 9004-02-8, Lipoprotein lipase 9004-06-2, Elastase 12629-01-5, Human growth hormone 56645-49-9, Cathepsin G 62031-54-3, FGF 83869-56-1, GM-CSF 127464-60-2, Vascular endothelial growth factor RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
                   (anti-heparanase monoclonal antibody for treating angiogenic conditions, tumors, inflammations, metastasis and autoimmune
                   diseases)
           ANSWER 8 OF 23 CAPLUS COPYRIGHT 2001 ACS SSION NUMBER: 2000:53938 CAPLUS
ACCESSION NUMBER:
 DOCUMENT NUMBER:
                                                              132:102821
                                                             Method of screening for potential anti-metastatic and
TITLE:
                                                              anti-inflammatory agents using mammalian heparanase as
                                                              a probe
                                                            a probe
Ben-Artzi, Hanna; Ayal-Hershkovitz, Maty; Vlodavsky,
Israel; Pecker, Iris; Peleg, Yoav; Miron, Daphna
Insight Strategy & Marketing Ltd., Israel; Hadasit
Medical Research Services & Development Ltd.;
Friedman, Mark M.
PCT Int. Appl., 70 pp.
CODEN: PIXXD2
INVENTOR (S):
 PATENT ASSIGNEE(S):
SOURCE:
 DOCUMENT TYPE:
                                                             Patent
 LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                              English
            PATENT NO.
                                                      KIND DATE
                                                                                                         APPLICATION NO. DATE
             WO 2000003036
                                                        Α1
                                                                   20000120
                                                                                                         WO 1999-US15643
                                                                                                                                                  19990712
                               003036 A1 20000120 w0 1999-01515643 19990/12
AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, ND, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TL, M, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
                    TM, TR, TT, UA, UG, US, US, VN, LS, MD, RU, TJ, TM
MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
6190875 B1 20010220 US 1998-113168 19980710
9948697 A1 20000201 AU 1999-48697 19990712
1007241 A1 20010509 EP 1999-932382 19990712
             US 6190875
             EP 1097241
                      R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO 2001000136 A 20010309 NO 2001-136 20010109
 NO 2001000136
PRIORITY APPLN. INFO.:
                                                                                                  US 1998-113168
US 1997-922170
                                                                                                                                           A 19980710
A2 19970902
                                                                                                  US 1998-109386 B2 19980702
WO 1999-US15643 W 19990712
AB Qual. and quant. methods are provided for testing an agent for its potential at inhibiting glycosidase catalytic activity, the methods including interacting a glycosidase enzyme with a glycosidase substrate in a presence of the agent and qual. or quant. evaluating an effect of the agent on the catalytic activity of the glycosidase enzyme toward the glycosidase substrate. Preferably the glycosidase enzyme is a heparanase enzyme and the glycosidase substrate is, resp., a heparanase substrate.

REFERENCE COUNT: 7
                                                              (2) Fuks; US 5362641 A 1994 CAPLUS (3) Gallo; US 5129877 A 1992 CAPLUS
  REFERENCE(S):
                                                             (4) Leshchiner; US 539351 A 1995 CAPLUS
(5) Lormeau; US 559351 A 1996 CAPLUS
(6) Nicholson; US 4859581 A 1999 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
             Antibodies
             RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (anti-heparanase; antimetastatic and antiinflammatory agent
                    screening with heparanase probe)
 L4 ANSWER 9 OF 23
ACCESSION NUMBER:
                                                           MEDLINE
                                                                                                                                          DUPLICATE 3
                                                 MEDLINE DUPLICATE 3
20010909022 MEDLINE
20476203 PubMed ID: 11021821
Expression of heparanase in normal, dysplastic, and neoplastic human colonic mucosa and stroma. Evidence for its role in colonic tumorigenesis.
Friedmann Y; Vlodavsky I; Aingorn H; Aviv A; Peretz T; Pecker I; Pappo O
Departments of Oncology and Pathology, Hadassah-Hebrew
 DOCUMENT NUMBER:
 AUTHOR:
 CORPORATE SOURCE:
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nd InSight Ltd., Rabin
                                                                            University Hospital, Jerusal
                                                                            Science Park, Rehovot, Israel.
AMERICAN JOURNAL OF PATHOLOGY, (200
Journal code: 3RS. ISSN: 0002-9440.
United States
                                                                                                                                                                                                 (2000 Oct) 157 (4) 1167-75.
SOURCE:
PUB. COUNTRY:
                                                                             Journal; Article; (JOURNAL ARTICLE)
LANGUAGE:
FILE SEGMENT:
                                                                             Abridged Index Medicus Journals; Priority Journals
ENTRY MONTH:
ENTRY DATE:
                                                                             200010
                                                                             Entered STN: 20010322
                 Y DATE: Entered STN: 20010322

Last Updated on STN: 20010322

Entered Medline: 20001025

The human heparanase gene, an endo-beta-glucuronidase that cleaves heparan sulfate at specific intrachain sites, has recently been cloned and shown to function in tumor progression and metastatic spread. Antisense digoxigenin-labeled heparanase RNA probe and monoclonal anti-human heparanase antibodies were used to examine the
                heparanase antibodies were used to examine the expression of the heparanase gene and protein in normal, dysplastic, and neoplastic human colonic mucosa. To our knowledge, this is the first systematic study of heparanase gene and protein in normal, dysplastic, and neoplastic human colonic mucosa. To our knowledge, this is the first systematic study of heparanase syression in human colon cancer. Both the heparanase gene and protein were expressed at early stages of neoplasia, already at the stage of adenoma, but were practically not detected in the adjacent normal-looking colon epithelium. Gradually increasing expression of heparanase was evident as the cells progressed from severe dysplasia through well-differentiated to poorly differentiated colon carcinoma. Deeply invading colon carcinoma cells showed the highest levels of the heparanase mRNA and protein associated with expression of both the gene and enzyme by adjacent desmoplastic stromal fibroblasts. A high expression was also found in colon carcinoma metastases to lung, liver, and lymph nodes, as well as in the accompanying stromal fibroblasts. Moreover, extracts derived from tumor tissue expressed much higher levels of the heparanase protein and activity as compared to the normal colon tissue. In all specimens, the heparanase gene and protein exhibited the same pattern of expression. These results suggest a role of heparanase in colon cancer progression and may have both prognostic and therapeutic applications.

. . . been cloned and shown to function in tumor progression and metastatic spread. Antisense digoxigenin-labeled heparanase RNA probe and monoclonal anti-human heparanase antibodies were used to examine the expression of the heparanase gene and protein in normal, dysplastic, and neoplastic human colonic mucosa. . . .
                    dysplastic, and neoplastic human colonic mucosa...
                   ANSWER 10 OF 23
                                                                                               MEDITAR
                                                                                                                                                                                                                     DUPLICATE 4
                                                                            2000229546 MEDLINE
20229546 PubMed ID: 10764835
Heparanase expression in invasive trophoblasts and acute
 ACCESSION NUMBER:
 DOCUMENT NUMBER:
                                                                              vascular damage.
                                                                             Dempsey L A; Plummer T B; Coombes S L; Platt J L
Department of Surgery, Mayo Clinic, Rochester, MN 55905,
 AUTHOR:
 CORPORATE SOURCE:
                                                                             GLYCOBIOLOGY, (2000 May) 10 (5) 467-75.
Journal code: BEL; 9104124. ISSN: 0959-6658.
                                                                             ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
 PUB. COUNTRY:
                                                                             English
Priority Journals
GENBANK-AF084467
  LANGUAGE .
 FILE SEGMENT:
OTHER SOURCE:
ENTRY MONTH:
                                                                            200005
Entered STN: 20000518
Last Updated on STN: 20000518
Entered Medline: 20000505
 ENTRY DATE:
                Last Updated on STN: 20000518
Entered Medline: 20000505

Heparan sulfate proteoglycans play a pivotal role in tissue function, development, inflammation, and immunity. We have identified a novel cDNA encoding human heparanase, an enzyme thought to cleave heparan sulfate in physiology and disease, and have located the HEP gene on human chromosome 4q21. Monoclonal antibodies against human heparanase located the enzyme along invasive extravillous trophoblasts of human placenta and along endothelial cells in organ xenografts targeted by hyperacute rejection, both sites of heparan sulfate digestion. Heparanase deposition was evident in arterial walls in normal tissues; however, vascular heparan sulfate cleavage was coincident with heparanase enzyme during inflammatory episodes. These findings suggest that heparanase elaboration and control of catalytic activity may contribute to the development and pathogenesis of vascular disease and suggest that heparanase intervention might be a useful therapeutic target.

. . . thought to cleave heparan sulfate in physiology and disease, and have located the HEP gene on human chromosome 4q21. Monoclonal antibodies against human heparanase located the enzyme along invasive extravillous trophoblasts of human placenta and along endothelial cells in organ xenografts targeted by hyperacute. . . .
 AB
                    ANSWER 11 OF 23 CAPLUS COPYRIGHT 2001 ACS SION NUMBER: 1999:723147 CAPLUS
  ACCESSION NUMBER:
  DOCUMENT NUMBER:
                                                                                                 131:332967
                                                                                                Genetically modified cells and methods for expressing recombinant heparanase and methods of purifying same Ben-Artzi, Hanna; Ayal-Hershkovitz, Maty; Yacoby-Zeevi, Oron; Pecker, Iris; Peleg, Yoav; Shlomi,
  INVENTOR(S):
                                                                                                 Yinon
  PATENT ASSIGNEE(S):
                                                                                                  Insight Strategy & Marketing Ltd., Israel; Friedman,
                                                                                                 Mark.
                                                                                                                    М.
  SOURCE:
                                                                                                PCT Int. Appl., 118 pp.
CODEN: PIXXD2
   DOCUMENT TYPE:
                                                                                                 Patent
                                                                                                 English
   LANGUAGE:
  LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                      PATENT NO.
                                                                                     KIND DATE
                                                                                                                                                                      APPLICATION NO. DATE
                                                                                                                                                                     WO 1999-US9256
                                                                                                                                                                                                                                     19990429
                      WO 9957244
                                                                                                           19991111
                                                                                        A1
                                                  A1 19991111 W0 1999-09255 19990429
AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
                                                  GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
                                   ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
9937705 A1 19991123 AU 1999-37705 19990429
1076689 A1 20010221 EP 1999-920135 19990429
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                     AU 9937705
EP 1076689
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NO 2000005100
                                                                                     20001228
                                                                                                                                                                                      20001010
 PRIORITY APPLN. INFO.:
                                                                                                                          US 1999-260038
                                                                                                                                                                                      19990302
                                                                                                                         WO 1999-US9256
                                                                                                                                                                                   19990429
                Bacterial, yeast and animal cells and methods for overexpressing
                 recombinant heparanase in cellular systems, methods of purifying recombinant heparanase therefrom and modified heparanase species which
               recombinant heparanase therefrom and modified heparanase species which serve as precursors for generating highly active heparanase by proteolysis. Heparanase is a glycosylated enzyme involved in catabolism of certain glycosaminoglycans, in tumor cell invasion and metastasis, and possibly in angiogenesis. It has potential therapeutic applications for viral infection, neurodegenerative diseases, restenosis, and atherosclerosis. A signal peptide was incorporated for effective protein secretion in yeast and bacteria and insect and mammalian cells. Protein secretion is achieved by induction by thrombin and calcium inonpohores and immune complexes and antigens and mitogens. This work describes prodn. of heparanase on a biotechnol. scale of at least half a liter growth medium by affinity purifn. This large scale propagation of animal cells is described in a Spinner-basket biotreactor. The heparanase enzyme is activated by digestion with a protease such as cathepsin L or trypsin at appropriate pH. A correctly folded catalytically active heparanase is generated.
                  generated.
   REFERENCE COUNT:
                                                                             (1) Insight Strategy & Marketing Ltd; WO 9911798 A1
 REFERENCE(S):
                                                                                         1999 CAPLUS
                                                                             (2) The Upjohn Co; WO 9504158 A1 1995 CAPLUS
                Antibodies
                ANLIBOOISE RE: ARG (Analytical reagent use); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (heparanase-specific; genetically modified cells and methods for expressing recombinant heparanase and methods of purifying same)
                ANSWER 12 OF 23 CAPLUS COPYRIGHT 2001 ACS SSION NUMBER: 1999:723067 CAPLUS
  ACCESSION NUMBER:
   DOCUMENT NUMBER:
                                                                              131:350261
  TITLE:
                                                                             Heparanase specific molecular probes and their use in
                                                                             research and medical applications
Pecker, Iris; Vlodavsky, Israel; Friedman, Yael;
Perets, Tuvia
  INVENTOR(S):
                                                                             Insight Strategy & Marketing Ltd., Israel; Hadasit Medical Research Services & Development Ltd.;
  PATENT ASSIGNEE(S):
                                                                             Friedman, Mark, M.
PCT Int. Appl., 90 pp.
CODEN: PIXXD2
  SOURCE:
   DOCUMENT TYPE:
                                                                              Patent
   LANGUAGE:
                                                                             English
  FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                 PATENT NO.
                                                                   KIND DATE
                                                                                                                                  APPLICATION NO. DATE
                           9957153 Al 19991111 WO 1999-US9255 19990429

W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MM, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

6177545 B1 20010123 US 1998-71739 19980501

8938706 A1 19991123 AU 1999-38706 19990429

1073682 A1 20010207 EP 1999-921513 19990429
                  WO 9957153
                  US 6177545
                  AU 9938706
                AU 9938/06 AI 1999II23 AU 1999-32170 19990429
EP 1073682 AI 20010207 EP 1999-921513 19990429
R: AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, NL, SE, IE, FI
NO 9906229 A 20000224 NO 1999-6229 19991215
RITY APPLN. INFO.: US 1998-71739 A 19980501
US 1997-922170 A2 19970902
   PRIORITY APPLN. INFO.:
US 1997-922170 A2 19970902

WO 1999-US9255 W 19990429

AB A variety of heparanase specific mol. probes which can be used for research and medical applications including diagnosis and therapy. Specific applications include the use of a heparanase specific mol. probe for detection of the presence, absence or level of heparanase expression; the use of a heparanase specific mol. probe for therapy of a condition assocd. with expression of heparanase; the use of a heparanase specific mol. probe for quantification of heparanase in a body fluid; the use of a heparanase specific mol. probe for targeted drug delivery; and the use of a heparanase specific mol. probe as a therapeutic agent.

REFERENCE COUNT: 14

REFERENCE (S): (2) Board Of Regents The University Of Tayon Succession
                                                                             WO 8801280 Al 1986 CAPPLUS

(3) Fuks; US 5362641 A 1994 CAPLUS

(4) Gewirtz; US 5618709 A 1997 CAPLUS

(6) Jin; Int J Cancer 1990, V45, Pl088 CAPLUS

(7) Kosir; J Surg Res 1997, V67, P98 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT
                  heparanase antibody nucleic acid probe cancer
                 neparanase antibody nucleic acid probe cancer
Primers (nucleic acid)
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(DNA; heparanase-specific antibodies and nucleic
acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
                              (polymerase chain reaction)
   ΙT
                          (RT-PCR (reverse transcription-PCR); heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
   IT
                  Lymphoma
                           (T-cell; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease,
                           diabetes and inflammation)
                  Ovary, neoplasm
                          (adenocarcinoma; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease,
                           diabetes and inflammation)
                            (agents; heparanase-specific antibodies and nucleic
                          acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
                  Proteins, general, processes
RL: REM (Removal or disposal); PROC (Process)
(body fluid; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease,
   IT
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diabetes and inflammation)
        diabetes and inframmation)
Mammary gland
(carcinoma, metastasis; heparanase-specific
antibodies and nucleic acid probes for diagnosis and therapy of
cancer, renal disease, diabetes and inflammation)
ΙT
         Bladder
         Mammary gland
         Prostate gland
(carcinoma; heparanase-specific antibodies and
nucleic acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
         Uterus, neoplasm
IT
               (cervix, metastasis; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
         Uterus, neoplasm
(cervix, squamous cell carcinoma; heparanase-specific
               antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
        Uterus, neoplasm
(cervix; heparanase-specific antibodies and nucleic
acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
ΙT
ΙT
         Intestine, neoplasm
         (colon, carcinoma, metastasis; heparanase-specific

(colon, carcinoma, metastasis; heparanase-specific
        colon, carcinoma, metastasis; heparanase-specific
antibodies and nucleic acid probes for diagnosis and therapy of
cancer, renal disease, diabetes and inflammation)
Intestine, neoplasm
(colon; heparanase-specific antibodies and nucleic
acid probes for diagnosis and therapy of cancer, renal disease,
ΙT
               diabetes and inflammation)
        Kidney, disease
(diabetic nephropathy; heparanase-specific antibodies
and nucleic acid probes for diagnosis and therapy of cancer, renal
disease, diabetes and inflammation)
IT
         Pleura (effusion; heparanase-specific antibodies and
1 T
               nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
         Animal cell
         Animal tissue
         Autoimmune disease
         Blood plasma
Body fluid
         Carcinoma
          DNA sequences
         Diabetes mellitus
         Drug delivery systems
Drug targeting
          Electrophoresis
         Hybridoma
          Inflammation
          Intestine, neoplasm
          Kidney, disease
          Labels
         Leukemia
          Liver, neoplasm
          Lymphoma
         Melanoma
         Multiple myeloma
         Nutriple myeloma
Neoplasm
Ovary, neoplasm
PCR (polymerase chain reaction)
Pancreas, neoplasm
Protein sequences
         Saliva
          Sepsis
         Skin, neoplasm
Stomach, neoplasm
          Urine
         Uterus, neoplasm
               (heparanase-specific antibodies and nucleic acid
               probes for diagnosis and therapy of cancer, renal disease, diabetes and
                inflammation)
         Nucleoside triphosphates
          mRNA
         MRNAM
RL: BSU (Biological study, unclassified); PUR (Purification or recovery);
THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
               (heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
         Antisense DNA
         Oligonucleotides
          RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
               (heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and
               inflammation)
         Antibodies
Probes (nucleic acid)
         RIO THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(heparanase-specific antibodies and nucleic acid
probes for diagnosis and therapy of cancer, renal disease, diabetes and
inflammation)
         Liver, neoplasm
               (hepatoma; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
        diabetes and inflammation)

Gene, animal

RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL

(Biological study); USES (Uses)

(hpa; heparanase-specific antibodies and nucleic
acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)

Diabetes mellitus

(insulindependent microslbyminuric benevance-specific
                (insulin-dependent, microalbuminuric; heparanase-specific
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antibodies and nucleic acid probes for
                                                                                                             nosis and therapy of
         cancer, renal disease, diabetes and inflammation) Diabetes mellitus
        (insulin-dependent, normoalbuminuric; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)

Mesothelium
ΙT
                 (mesothelioma; heparanase-specific antibodies and
                nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
          Intestine, neoplasm
Liver, neoplasm
          Neoplasm
          Ovary, neoplasm
Pancreas, neoplasm
Prostate gland
          Skin, neoplasm
Stomach, neoplasm
Uterus, neoplasm
                (metastasis; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
         Antigens
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(moiety; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease,
                 diabetes and inflammation)
          Antibodies
          RR: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(monoclonal; heparanase-specific antibodies and
nucleic acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
          Bladder
          Mammary gland
                 (neoplasm, metastasis; heparanase-specific antibodies
                and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
          Bladder
          Mammary gland
          Prostate gland
Prostate gland
(neoplasm; heparanase-specific antibodies and
nucleic acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
         Kidney, disease
(nephritis, hemorrhagic; heparanase-specific
antibodies and nucleic acid probes for diagnosis and therapy of
cancer, renal disease, diabetes and inflammation)
          Kidney, disease
                 (nephrotic syndrome; heparanase-specific antibodies
                and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
ΙT
          RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (primer; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease,
         acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
Nucleic acids
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(probe; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
ΙT
          Separation
                 (size; heparanase-specific antibodies and nucleic
                acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
IT
          Neoplasm
                (solid; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
           Drug delivery systems
                 (solns:) heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
IT
          Carcinoma
                (squamous cell; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
          Carcinoma
                (teratocarcinoma; heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
          Hematopoietic precursor cell
(tumors; heparanase-specific antibodies and nucleic
acid probes for diagnosis and therapy of cancer, renal disease,
diabetes and inflammation)
          221113-49-1, Heparanase (human gene hpa) RL: PRP (Properties)
          RL: PRP (Properties)

(amino acid sequence: heparanase-specific antibodies
and nucleic acid probes for diagnosis and therapy of cancer, renal
disease, diabetes and inflammation)

89800-66-8, Heparanase

RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(heparanase-specific antibodies and nucleic acid
probes for diagnosis and therapy of cancer, renal disease, diabetes and
           inflammation)
249927-19-3 249927-20-6 249927-21-7 249927-22-8
RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES
                 (heparanase-specific antibodies and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and
                  inflammation)
           250215-44-2
           RL: PRP (Properties)
(nucleotide sequence; heparanase-specific antibodies
          and nucleic acid probes for diagnosis and therapy of cancer, renal disease, diabetes and inflammation)
9012-90-2, DNA polymerase
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
                  (thermostable; heparanase-specific antibodies and
```

nucleic acid probes for diagnosis and diabetes and inflammation)

L4 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:566196 CAPLUS

DOCUMENT NUMBER:

131:181667

Human platelet heparanase polypeptides, polynucleotide molecules that encode them, and methods for the identification of compounds that alter heparanase

INVENTOR (S):

Heinrikson, Robert L.; Fairbanks, Michael B.; Mildner,

Pharmacia & Upjohn Company, USA PATENT ASSIGNEE (S): PCT Int. Appl., 57 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 9943830 TJ, TM

RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

9927591 Al 19990915 AU 1999-27591 19990218

1060252 A2 20001220 EP 1999-908073 19990218 AU 9927591 EP 1060252

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
PRIORITY APPLN. INFO.: US 1998-75706 P 19980224

US 1998-79401 WO 1999-US1489 P 19980326 W 19990218

WO 1999-US1489 W 19990218

The present invention provides isolated human heparanase polypeptides, and the isolated polynucleotide mols. that encode them, as well as vectors and host cells comprising such polynucleotide mols. Heparanase is purified from human platelets (20.mu.g from 2000 mL platelet-rich plasma) by heparin-Sepharose chromatog., size exclusion chromatog. on Superdex-75, and heparin HiTrap column chromatog. The heparanase transcript encodes a 530-amino acid residue prepro enzyme which is processed to form 8-kDa and 56-kDa subunits. The invention also provides a method for the identification of an agent that alters heparanase activity.

Antibodies

Antibodies

RL: ARG (Analytical reagent use); BPN (Biosynthetic preparation); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES

(human platelet heparanase polypeptides, polynucleotide mols. that encode them, and methods for the identification of compds. that alter heparanase activity)

L4 ANSWER 14 OF 23 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:511264 CAPLUS

DOCUMENT NUMBER:

131:141481

Human heparanase obtained from SV-40-transformed cell line, its cDNA and amino acid sequences, recombinant expression and its biological, diagnostic and therapeutic uses

INVENTOR(S): PATENT ASSIGNEE(S):

therapeutic uses
Nakajima, Motowo; Toyoshima, Minako
Novartis A.-G., Switz.; Novartis-Erfindungen
Verwaltungsgesellschaft m.b.H.
PCT Int. Appl., 40 pp.
CODEN: PIXXD2

SOURCE:

DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PA: | TENT | NO. |     | KI  | ND  | DATE |      |     |     |      |      | ON N |     | DATE |      |     |     |
|-----|------|-----|-----|-----|-----|------|------|-----|-----|------|------|------|-----|------|------|-----|-----|
| WO  | 9940 | 207 |     | A   | 1   | 1999 | 0812 |     |     |      |      |      |     | 1999 | 0205 |     |     |
|     | W:   | AL, | AM, | AT, | AU, | AZ,  | BA,  | BB, | BG, | BR,  | BY,  | CA,  | CH, | CN,  | CU,  | CZ, | DE, |
|     |      |     |     |     |     |      |      |     |     |      |      |      |     | IL,  |      |     |     |
|     |      | KE, | KG, | KP, | KR, | KZ,  | LC,  | LK, | LR, | LS,  | LT,  | LU,  | LV, | MD,  | MG,  | MK, | MN. |
|     |      | MW, | MX, | NO, | NZ, | PL,  | PT,  | RO, | RU, | SD,  | SE,  | SG,  | SI, | SK,  | SL,  | ΤJ, | TM, |
|     |      | TR, | TT, | UA, | UG, | US,  | UΖ,  | VN, | YU, | ZW,  | AM,  | AZ,  | BY, | KG,  | ΚZ,  | MD, | RU, |
|     |      | TJ, | TM  |     |     |      |      |     |     |      |      |      |     |      |      |     |     |
|     | RW:  | GH, | GM, | ΚE, | LS, | MW,  | SD,  | SZ, | UG, | ZW,  | AT,  | ΒE,  | CH, | CY,  | DE,  | DK, | ES, |
|     |      | FI, | FR, | GB, | GR, | ΙE,  | IT,  | LU, | MC, | NL,  | PT,  | SE,  | BF, | ВJ,  | CF,  | CG, | CI, |
|     |      | CM, | GΑ, | GN, | GW, | ML,  | MR,  | ΝE, | SN, | TD,  | TG   |      |     |      |      |     |     |
| ΑU  | 9928 | 319 |     | A   | 1   | 1999 | 0823 |     | A.  | U 19 | 99-2 | 8319 |     | 1999 | 0205 |     |     |
| EP  | 1054 | 980 |     | А   | 1   | 2000 | 1129 |     | E   | P 19 | 99-9 | 0885 | 4   | 1999 | 0205 |     |     |
|     | R:   | ΑT, | BE, | CH, | DE, | DK,  | ES,  | FR, | GB, | GR,  | ΙT,  | LI,  | LU, | NL,  | SE,  | MC, | PT, |
|     |      | IE, | FI  |     |     |      |      |     |     |      |      |      |     |      |      |     |     |
|     |      |     |     |     |     |      |      |     |     |      |      |      |     |      |      |     |     |

PRIORITY APPLN. INFO .:

to 20 kDa fragments. REFERENCE COUNT:

REFERENCE (S):

(1) Christopher, R; WO 9633726 A 1996 CAPLUS

(2) Goshen; Molecular Human Reproduction 1996, V2(9), P679 CAPLUS

(3) Hadassah Med Org; WO 9102977 A 1991 CAPLUS

(4) Hamdorf, B; WO 992

(6) Hoogwerf, A; WO 9504158 A 1995 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
This invention presents a polynuclectide (cDNA mol.) encoding human
heparanase, which was obtained from the SV-40-transformed fibroblast cell
line ATCC CCL 75.1. The invention also provides a hybrid vector contg.
the human heparanase polynuclectide, and a host cell transformed with the
vector. The invention further provides an antibody that specifically
recognizes and binds the heparanase, and use of this
antibody in treatment of, diseases assocd. with abnormal expression
or activity of heparanase. Still further, the invention presents the
therapeutic, diagnostic and biol. uses of human heparanase including the
use of heparanase to identify agonist or antagonists of heparanase.
Finally, the invention presents an oligonuclectide capable of specifically
hybridizing with the human heparanase polynuclectide, and use of the human
heparanase, olynuclectide in gene therapy. The CDNA sequence encoding
human heparanase, as well as the amino acid sequences encoding the prepro
and mature forms of heparanase are claimed. The heparanase is an
endoglucuronidase capable of specifically degrading heparan sulfate into 6
to 20 kDa fragments. to 20 kDa fragments. RRL: ARG (Analytical reagent use); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (anti-human heparanase, used in treatment of diseases; human heparanase obtained from SV-40-transformed cell line, its cDNA and heparanase, used in treatment of diseases; human heparanase obtained from SV-40-transformed cell line, its cDNA and amino acid sequences, recombinant expression and its biol., diagnostic and therapeutic uses) ANSWER 15 OF 23 CAPLUS COPYRIGHT 2001 ACS SSION NUMBER: 1999:303252 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 130:322335 Isolated nucleic acid molecule encoding mammalian endoglucuronidase and its therapeutic uses for enhancing wound healing and related disorders Freeman, Craig Geoffrey; Hulett, Mark Darren; Parish, Christopher Richard; Hamdorf, Brenton James The Australian National University, Australia INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 112 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE WO 9921975 A1 19990506 9921975 A1 19990506 W0 1998-AU898. 19981028
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
9910109 A1 19990517 AU 1998-10109 19981028
9813296 A 20000822 BR 1998-13296 19981028 AU 9910109 ZA 9809824 A 19971028 A 19971209 PRIORITY APPLN. INFO.: AU 1997-62 AU 1997-812

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US 1998-181336
                    A 19981028
W 19981028
WO 1998-AU898
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The present invention relates to isolated or recombinant mammalian endoglucuronidase enzymes, polypeptides and peptides, in particular human, murine and rat heparanases, and genetic sequences encoding the enzymes. The full-length cDNA sequence of the human heparanase enzyme contains an open reading frame encoding a 543-amino acid protein. Also included are uses in the detn. and characterization of chem. compds., proteins, polypeptides, small mols. and macromols. capable of inhibiting metastasis, angiogenesis, angioplasty-induced restenosis, atherosclerosis, inflammation, promote wound healing and otherwise modulate physiol. processes involving heparanase cleavage of heparan sulfate. The invention further relates to a method of altering, modifying or otherwise modulating the level of expression of mammalian heparanase in a cell. A further aspect of the invention relates to immunoreactive mols. capable of binding to and/or inhibiting mammalian heparanase, in particular The present invention relates to isolated or recombinant mammalian AB monoclonal antibodies. A still further aspect of the invention contemplates the use of heparanase as an agent to promote the processes of wound healing.
REFERENCE COUNT:

REFERENCE(S):

RENCE COUNT: 2

RENCE(S): (1) Turnbull, J; Biochem J V265, P715 CAPLUS

(2) Turnbull, J; Biochem J 1991, V273, P553 CAPLUS

The present invention relates to isolated or recombinant mammalian
endoglucuronidase enzymes, polypeptides and peptides, in particular human,
murine and rat heparanases, and genetic sequences encoding the enzymes.

The full-length cDNA sequence of the human heparanase enzyme contains an
open reading frame encoding a 543-amino acid protein. Also included are
uses in the detn. and characterization of chem. compds., proteins,
polypeptides, small mols. and macromols. capable of inhibiting metastasis,
angiogenesis, angioplasty-induced restenosis, atherosclerosis,
inflammation, promote wound healing and otherwise modulate physiol.
processes involving heparanase cleavage of heparan sulfate. The invention
further relates to a method of altering, modifying or otherwise modulating
the level of expression of mammalian heparanase in a cell. A further
aspect of the invention relates to immunoreactive mols. capable of binding aspect of the invention relates to immunoreactive mols. capable of binding to and/or inhibiting mammalian heparanase, in particular monoclonal antibodies. A still further aspect of the invention contemplates the use of heparanase as an agent to promote the processes of wound healing.

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MEDLINE DUPLICATE 5
1999386667 MEDLINE
99386667 PubMed ID: 10455023
Evidence that platelet and tumour heparanases are similar
L4 ANSWER 16 OF 23 ACCESSION NUMBER:
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DOCUMENT NUMBER:

AUTHOR

enzymes.
Freeman C; Browne A M; Parish C R
Division of Immunology and Cell Biology, John Curtin School
of Medical Research, Australian National University, CORPORATE SOURCE:

Canberra, ACT, 2601, Austral Craig.Freeman BIOCHEMICAL JOURNAL, (1999 Sep 1) 342 ( Pt 2) JOURNAL CODE: 090; 2984726R. ISSN: 0264-6021. ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) Craig.Freeman@anu.edu.au SOURCE:

PUB. COUNTRY:

FILE SEGMENT: Priority Journals ENTRY MONTH: 199911

Entered STN: 20000111 ENTRY DATE:

IN MONTH: 19961
If DATE: Entered STN: 20000111
Last Updated on STN: 20000111
Entered Medline: 19991101
In order to enter tissues, blood-borne metastatic tumour cells and leucocytes need to extravasate through the vascular basal lamina (BL), a process which involves a battery of degradative enzymes. A key degradative enzyme is the endoglycosidase heparanase, which cleaves heparan sulphate (HS), an important structural component of the vascular BL. Previously, tumour-derived heparanase activity (which has been shown to be related to the metastatic potential of murine and human melanoma cell lines) was reported to cleave HS and be inhibited by heparin, as distinct from human platelet heparanase, which cleaved both substrates (Nakajima, Irimura and Nicolson (1938) J. Cell Biochem. 36, 157-167]. We recently reported the purification of human platelet heparanase and showed that the enzyme is a 50-kDa endoglucuronidase [Freeman and Parish (1998) Biochem. J. 330, 1341-1350]. We now report the purification and characterization of heparanase activity from highly metastatic rat 13762 MAT mammary adenocarcinoma and human HCT 116 colonic carcinoma cells and from rat liver using essentially the same procedure that was reported for liver using essentially the same procedure that was reported for purification of the human platelet enzyme. The rat 13762 MAT tumour enzyme, which has a native M(r) of 45 kDa when analysed by gel-filtration chromatography and by SDS/PAGE, was observed to be an endoglucuronidase that degraded heparin and HS to fragments of the same sizes as the human chromatography and by SDS/PAGE, was observed to be an endoglotation as that degraded heparin and HS to fragments of the same sizes as the human platelet enzyme does. N-deglycosylation of both the human platelet and rat 13762 MAT tumour enzymes gave, in each case, a 41-kDa band by SDS/PAGE analysis, demonstrating that the observed difference in M(r) between the platelet and tumour enzymes may have been due largely to differences in the relative amounts of N-glycosylation. Two peptides were isolated following Endoproteinase Lys-C digestion of both the human platelet and rat 13762 MAT tumour heparanases and were shown to be highly similar. Both the rat liver and human colonic carcinoma heparanases also degraded both heparin and HS to fragments of the same sizes as the human platelet enzyme does. Western-blot analysis of an SDS/PAGE gel using antibodies raised against human platelet heparanase demonstrated that human platelet, human tumour and rat tumour heparanases were immunochemically cross-reactive. In conclusion, because of the similarities in their sizes, substrate specificities, peptide sequences and immunoreactivities, we propose that heparanase activities present in human platelets, rat liver and in rat and human tumour cells are, in fact, mediated by a similar enzyme.

encyme. . . . HS to fragments of the same sizes as the human platelet enzyme does. Western-blot analysis of an SDS/PAGE gel using antibodies raised against human platelet heparanase demonstrated that human platelet, human tumour and rat tumour heparanases were immunochemically cross-reactive. In conclusion, because of the similarities in.

ANSWER 17 OF 23 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER:

1999108041 MEDLINE 99108041 PubMed ID: 9889056

Degradation of basement membrane by prostate tumor

heparanase. Kosir M A; Wang W; Zukowski K L; Tromp G; Barber J AUTHOR: CORPORATE SOURCE:

SOURCE:

Kosir M A; Wang W; Zukowski K L; Fromp G; Barber J Department of Surgery, Wayne State University School of Medicine, Detroit, Michigan, 49201, USA. KOSIR MARY ANNEDETROIT.VA.GOV JOURNAL OF SURGICAL RESEARCH, (1999 Jan) 81 (1) 42-7. Journal code: K7B; 0376340. ISSN: 0022-4804.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE: FILE SEGMENT: Priority Journals

ENTRY MONTH: 199902

ENTRY DATE:

MONTH: 199902

( DATE: Entered STN: 19990311

Last Updated on STN: 19990311

Entered Medline: 19990223

BACKGROUND: The degradation of basement membrane (BM) by cancer is an

Entered Medline: 19990213
BACKGROUND: The degradation of basement membrane (BM) by cancer is an important event that characterizes invasive biological behavior. A component of BM is heparan sulfate proteoglycan (HSPG). The glycanase(s) that degrade HSPG in EM are not yet isolated. We recently identified HSPG-degrading activity (PC-3M heparanase) in the conditioned media (CM) of malignant prostate carcinoma cells (PC-3M and LNCaP C4-2).

Antibodies (Abs) to a recently isolated heparanase from human platelets (CTAP-III), cross-react with PC-3M heparanase although they differ in size; under reduced conditions PC-3M heparanase is 60 kDa whereas CTAP-III is 10 kDa by polyacrylamide gel electrophoresis. PC-3M heparanase therefore shares homology with CTAP-III. The purpose of this study was to test the inhibition of PC-3M heparanase by Abs specific to the N- and C-terminals of CTAP-III. MATERIALS AND METHODS: CM from PC-3M and LNCaP C4-2 cells were tested for heparanase activity. Each reaction contained substrate as [3H]glucosamine-labeled HSPG (>50 kDa) from the BM of the EHS tumor, CM from PC-3M or LNCaP C4-2 cells, and inhibitor or buffer (negative control). Protease inhibitors were present throughout. After incubation for 3-20 h at 37 degreesC and pH 5.8, the reaction was stopped with 0.2% SDS. Each reaction mixture was centrifuged in an Ultrafree-MC 30,000 NMWL filter unit (Millipore) and radioactivity in the filtrate counted by scintillation counting. Results. For both cell lines, there was a linear relationship between the amount (microgram) of CM and degradation of HSPG. Degradation was inhibited by 54.1% (mean) using carrageenan lambda (10 microgram/ml), a nonspecific glycanase inhibitor (P < 0.05 by ANOVA). Ab to the N-terminus of CTAP-III (anti-Hep A) reduced degradation by 10-50% (mean 31.1%) and to the C-terminus (anti-Hep C) by 38.8-64.3% (mean 51.1%) (P < 0.003 by ANOVA). CONCLUSIONS: The degradation of HSPG by malignant prostate cancer cell lines is inhibited by both a nonspecific glycanase inhibitor,

heparanase is 60.

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR:

SOURCE:

1998008076 MEDLINE 98008076 PubMed ID: 9581574 Major co-localization of the extracellular-matrix

degradative enzymes heparanase and gelatinase in tertiary granules of human neutrophils.

Mollinedo F; Nakajima M; Llorens A; Barbosa E; Callejo S; Gajate C; Fabra A

CORPORATE SOURCE:

Gajate C; Fabra A
Laboratory of Signal Transduction and Leucocyte Biology,
Instituto de Biologia y Genetica Molecular, Facultad de
Medicina, Consejo Superior de Investigaciones
Cientificas-Universidad de Valladolid, C/ Ramon y Cajal,
E-47005 Valladolid, Spain.
BIOCHEMICAL JOURNAL, (1997 Nov 1) 327 ( Pt 3) 917-23.
JOURNAL CODE: 9YO; 2984726R. ISSN: 0264-6021.

PUB. COUNTRY:

ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) LANGUAGE:

English Priority Journals

FILE SEGMENT: ENTRY MONTH: 199805 ENTRY DATE:

SEGMENT: Priority Journals

YMONTH: 199805

INTO ATE: Entered STN: 19980529

Last Updated on STN: 20000303

Entered Medline: 19980520

The expression of cell-surface adhesion proteins and the release of extracellular-matrix degradative enzymes constitute crucial processes for the attachment of neutrophils to the endothelium and for the subsequent extravasation of these cells through the endothelial layer. We have analysed in resting human neutrophils the subcellular localization of heparanase, a heparan-sulphate-degrading endoglycosidase that can degrade basement-membrane components, thereby facilitating neutrophil passage into the tissue during an inflammatory reaction. By subcellular fractionation of postnuclear supernatants from resting human neutrophils on continuous sucrose gradients, we have found that heparanase activity was mainly located in gelatinase-containing tertiary granules. Using a specific antibody, the 96-kDa heparanase protein was further located in the gelatinase-rich subcellular fractions. Following immunoblotting and immunoprecipitation analysis in the distinct subcellular fractions, we also found co-localization of heparanase and Mol (CDIb/CDI8), a leucocyte integrin involved in the attachment of neutrophils to the endothelium, in the fractions enriched in gelatinase-containing tertiary granules. Treatment of human neutrophils with tumour necrosis factor or granulocyte/macrophage colony-stimulating factor induced an increase in the CDIb/CDI8 cell-surface expression, as well as the release of both gelatinase (matrix metalloproteinase-9) and heparanase, but not of other granule markers, indicating a major co-localization of gelatinase, heparanase and CDIbb/CDI8 in the same organelle. Furthermore, confocal laser scanning microscopy using specific antibodies against gelatinase and heparanase and CDIbb/CDI8 in the gelatinase-containing tertiary granule supports the notion that mobilization of this organelle can regulate extravasation of human neutrophils.

On continuous sucrose gradients, we h neutrophils.

neutrophils.
. . . on continuous sucrose gradients, we have found that heparanase activity was mainly located in gelatinase-containing tertiary granules. Using a specific antibody, the 96-kDa heparanase protein was further located in the gelatinase-rich subcellular fractions. Following immunoblotting and immunoprecipitation analysis in the distinct subcellular fractions, we. . indicating a major co-localization of gelatinase, heparanase and CD11b/CD18 in the same organelle. Furthermore, confocal laser scanning microscopy using specific antibodies against gelatinase and heparanase revealed a major co-localization of both enzymes in intracellular cytoplasmic granules. The major localization of beparanase and CD11b/CD18 in the. major localization of heparanase and CD11b/CD18 in the. .

DUPLICATE 8 MEDLINE ANSWER 19 OF 23

ACCESSION NUMBER: DOCUMENT NUMBER:

97223306 97223306 MEDLINE PubMed ID: 9070190

Human prostate carcinoma cells produce extracellular

AUTHOR:

Kosir M A; Ouinn C C; Zukowski K L; Grignon D J; Ledbetter

Surgical Service, VA Medical Center, Detroit, Michigan CORPORATE SOURCE:

Ag201, USA.
JOURNAL OF SURGICAL RESEARCH, (1997 Jan) 67 (1) 98-105.
JOURNAL code: K7B; 0376340. ISSN: 0022-4804.
United States SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

English LANGUAGE: FILE SEGMENT:

Priority Journals 199704

ENTRY MONTH: ENTRY DATE:

human prostate carcinoma cells show heparanase activity in conditioned medium that degrades heparin and BM HSPG and is detected by antibody to platelet heparanase. In addition, the membrane-associated staining in tissue sections of prostate cancer strongly correlates with the biochemical and immunological detection in

conditioned medium of human PC-3M cells. . . . human prostate carcinoma cells show neparanase activity in conditioned medium that degrades heparin and BM HSPG and is detected by antibody to platelet heparanase. In addition, the membrane-associated staining in tissue sections of prostate cancer strongly correlates with the biochemical and immunological detection in. 3 MEDLINE DUPLICATE 9
94148528 MEDLINE
94148528 PubMed ID: 8314313
Immunoselection of GRP94/endoplasmin from a KNRK
cell-specific lambda gtl1 library using antibodies
directed against a putative heparanase
amino-terminal peptide.
De Vouge M W; Yamazaki A; Bennett S A; Chen J H; Shwed P S;
Couture C; Birnboim H C
Ottawa Regional Cancer Centre, Ontario, Canada. L4 ANSWER 20 OF 23 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: AUTHOR: Ottawa Regional Cancer Centre, Ontario, Canada. INTERNATIONAL JOURNAL OF CANCER, (1994 Jan 15) 56 (2) CORPORATE SOURCE: SOURCE: 286-94. Journal code: GQU; 0042124. ISSN: 0020-7136. PUB. COUNTRY: United States Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: OTHER SOURCE: ENTRY MONTH: Priority Journals GENBANK-S69315; GENBANK-S69316 199403 If DATE: Entered STN: 19940330

Last Updated on STN: 19960129

Entered Medline: 19940322

Induction of an invasive phenotype by metastatic tumour cells results in part from inappropriate expression of extracellular matrix-degrading enzymes normally involved in embryonic morphogenesis, tissue remodelling, angiogenesis and wound healing. Such enzymes include endoglycosidases that degrade heparan sulfate (HS) in endothelial basement membrane, as well as better characterized proteases. Heparanase, an endo-beta-D-glucuronidase initially detected in B16 melanoma cells, has been described as a M(r) 96,000 glycoprotein with p1 of 5.2, and has been immunolocalized to the cell surface and cytoplasm. We have utilized a polyacrylamide-gel-based HS degradation assay to demonstrate that KNRK, a rat kidney fibroblast cell line transformed by v-K-ras, exhibits HS-degrading activity similar to that of B16F10 mouse melanoma cells. To immunoselect heparanase-expressing clones from a KNRK-cell-specific lambda gtl1 cDNA library, we have also prepared a rabbit anti-serum directed against a putative amino-terminal ENTRY DATE: Entered STN: 19940330 clones from a KNRK-cell-specific lambda gtll cDNA library, we have also prepared a rabbit anti-serum directed against a putative amino-terminal peptide of B16F10 cellular heparanase. Lysogens from one clone expressed a beta-galactosidase fusion protein whose staining with peptide anti-serum was inhibited by competition with excess peptide. Dideoxy-mediated sequencing of the insert termini of this recombinant revealed that it represents a rat homologue of M(r) 94,000 glucose-regulated protein (GRP94/endoplasmin), a molecular chaperone that contains the exact amino-terminal sequence previously attributed to heparanase. Our results call into question the specificity of this peptide sequence, as well as previous immunolocalization studies of heparanase carried out using such anti-sera. Immunoselection of GRP94/endoplasmin from a KNRK cell-specific lambda gtl1 library using antibodies directed against a putative heparanase amino-terminal peptide. L4 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1992:190176 CAPLUS 116:190176 DOCUMENT NUMBER: TITLE: Antibodies, kits, and methods for immunochemical localization of heparanase in mouse and human melanomas, and characterization of melanoma heparanase Nicolson, Garth L.; Nakajima, Motowo; Jin, Li INVENTOR(S): University of Texas System, USA PCT Int. Appl., 82 pp. CODEN: PIXXD2 PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. PATENT NO. KIND DATE DATE WO 9119197 A1 19911212 WO 1991-US3832 19910530 W: AU, CA, JP
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE
9182317 A1 19911231 AU 1991-82317 19910530 W: AU, C.,
RW: AT, BE, CH, DE, DK, ES,
AU 9182317
A1 19911231
AU 641269
B2 19930916
EP 532695
A1 19930324
EP 1991-913555
19910530
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE
JP 05509403
T2 19931222
JP 1991-512410
19910530
W0 1991-US3832
19910530
W0 1991-US3832
19910530

Andoglycosidase (esp. PRIORITY APPLN. INFO.: Antibodies to a glycosaminoglycan endoglycosidase (esp. heparanase), as well as kits and methods employing the antibodies, are disclosed. Antibodies against an N-terminal heparanase peptide are produced. These antibodies are used for the detection of heparan sulfate endoglycosidase in human and murine tumors. Purifn. of melanoma heparanase is described. A hemocyanin-coupled heparanase-derived peptide was used as an immunogen for antibody prodn. Also described is prepn. and reactivity of various substrates (e.g. desulfated or desulfated and acetylated heparan sulfate) with melanoma heparanase. The anti-heparanase antibodies of the invention stained metastatic melanoma cells, but did not stain surrounding tissue.

Antibodies to a glycosaminoglycan endoglycosidase (esp. heparanase), as well as kits and methods employing the antibodies, are disclosed. Antibodies against an N-terminal heparanase peptide are produced. These antibodies are used for the detection of heparan sulfate endoglycosidase in human and murine tumors. Purifn. of melanoma heparanase is described. A hemocyanin-coupled heparanase-derived peptide was used as an immunogen for antibody prodn. Also described is prepn. and reactivity of various substrates (e.g. desulfated or desulfated and acetylated heparan sulfate) with melanoma heparanase. The anti-heparanase melanoma cells, but did not stain surrounding tissue. did not stain surrounding tissue. glycosaminoglycan endoglycosidase antibody; heparanase melanoma ST antibody Animal cell line (B16-F10, heparanase purifn. from, antibodies for

```
melanoma localization in relation to)
IΤ
              Melanoma
                        (heparanase of, purifn. of and antibodies for localization of)
ΙT
               Antibodies
               RL: ANST (Analytical study)
              (to heparanase, melanoma localization in relation to)
Hemocyanins
Proteins, specific or class
RL: ANST (Analytical study)
(conjugates, with heparanase peptide, for anti-
heparanase antibody prodn.)
Melanoma
ΙT
                        (metastatic, anti-heparanase antibodies staining
               89800-66-8, Heparanase
                                                                                           140879-15-8
              RE: ANST (Analytical study)
(antibodies to and immunochem. detection of, melanoma localization in relation to)
139775-54-5 139878-83-4
               RL: ANST (Analytical study)
                         (for anti-heparanase antibody prodn.)
                                                                                                                                                                           DUPLICATE 10
               ANSWER 22 OF 23
                                                                           MEDLINE
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                            90277246 MEDLINE
90277246 PubMed ID: 2351486
                                                              Immunochemical localization of heparanase in mouse and human melanomas.
                                                              Jin L: Nakajima M; Nicolson G L
AUTHOR:
                                                             Department of Tumor Biology, University of Texas M.D. Anderson Cancer Center, Houston 77030.
CORPORATE SOURCE:
                                                             P30-CA16672 (NCI)
R01-CA41524 (NCI)
R35-CA44352 (NCI)
CONTRACT NUMBER:
                                                               INTERNATIONAL JOURNAL OF CANCER, (1990 Jun 15) 45 (6)
SOURCE:
                                                              1088-95.
                                                             Journal code: GQU; 0042124. ISSN: 0020-7136. United States
PUB. COUNTRY:
                                                              Journal; Article; (JOURNAL ARTICLE)
 LANGUAGE:
                                                              English
FILE SEGMENT:
ENTRY MONTH:
                                                             Priority Journals
                                                              199007
                                                              Entered STN: 19900824
ENTRY DATE:
                                                             Last Updated on STN: 19970203
Entered Medline: 19900719
             Entered Medline: 19900719
Heparanase, an endo-beta-D-glucuronidase, has been associated with melanoma metastasis. Polyclonal antibodies directed against the murine N-terminal heparanase peptide detected a Mr approximately 97,000 protein on SDS-PAGE of mouse melanoma and human melanoma cell lysates. In an indirect immunocytochemical study, human A375-SM and mouse B 16-BL6 melanoma cells were stained with the anti-heparanase antibodies. Heparanase antigen was localized in the cytoplasm of permeabilized melanoma cells as well as at the cell surface of unpermeabilized cells. Immunohistochemical staining of frozen sections from syngenic mouse lungs containing micrometastases of B16-BL6 melanoma demonstrated heparanase localized in metastatic melanoma cells. Similar
              from syngeneic mouse lungs containing micrometastases of B16-BL6 melanoma demonstrated heparanase localized in metastatic melanoma cells. Similar studies using frozen sections of malignant melanomas resected from patients indicated that heparanase is localized in invading melanoma cells. Our studies suggest that (a) the N-terminus of the heparanase molecule in mouse and human is antigenically related; (b) heparanase antigens are localized at the cell surface and in the cytoplasm of metastatic human and mouse melanoma cells; and (c) heparanase antigens are enriched in invasive and metastatic murine and human melanomas in vivo.

. . . melanoma cell lysates. In an indirect immunocytochemical study, human A375-SM and mouse B 16-BL6 melanoma cells were stained with the anti-heparanase antipodies. Heparanase
                anti-heparanase antibodies. Heparanase antigen was localized in the cytoplasm of permeabilized melanoma cells as well as at the cell surface of unpermeabilized cells. . . .
                                                                            MEDLINE
               ANSWER 23 OF 23
 ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                              87109488
87109488
                                                                                                    MEDITNE
                                                                                             PubMed ID: 2433294
                                                              Soluble antigen induces T lymphocytes to secrete an endoglycosidase that degrades the heparan sulfate moiety of subendothelial extracellular matrix.
 TITLE:
 AUTHOR:
                                                              Fridman R; Lider O; Naparstek Y; Fuks Z; Vlodavsky I; Cohen
                                                              I K
CA 30289 (NCI)
NS 1868 (NINDS)
JOURNAL OF CELLULAR PHYSIOLOGY, (1987 Jan) 130 (1) 85-92.
JOURNAL code: HNB; 0050222. ISSN: 0021-9541.
United States
 CONTRACT NUMBER:
 SOURCE:
 PUB. COUNTRY:
                                                               Journal; Article; (JOURNAL ARTICLE)
  LANGUAGE:
                                                               English
                                                              Priority Journals
  FILE SEGMENT:
                                                               198703
              Y MONTH: 198703
Y DATE: Entered STN: 19900303
Last Updated on STN: 19970203
Entered Medline: 19870311
The antigen-mediated induction of heparanase, an endoglycosidase capable of degrading heparan sulfate from the subendothelial extracellular matrix (ECM), was investigated in a rat T lymphocyte cell line reactive against the basic protein (BP) of myelin. We found that nonactivated T lymphocytes could be induced to express heparanase activity following exposure to soluble but not to ECM-bound BP. The induction of heparanase was immunologically specific and independent of the presence of syngeneic or allogeneic antigen presenting cells (APC). However, anti-TA antibodies inhibited heparanase expression. Soluble BP induced secretion of heparanase into the culture medium within minutes, despite inhibition of protein synthesis. Cell lysates of T lymphocytes contained heparanase activity. Thus, T lymphocytes secrete a preformed heparanase following exposure to specific antigen.

. . . of heparanase was immunologically specific and independent of the presence of syngeneic or allogeneic antigen presenting cells (APC). However, anti-TA antibodies inhibited heparanase expression. Soluble BP induced secretion of heparanase into the culture medium within minutes, despite inhibition of protein synthesis. Cell lysates. . .
                                                              Entered STN: 19900303
 ENTRY DATE:
```

COST IN U.S. DOLLARS

FULL ESTIMATED COST

COST IN U.S. DOLLARS

ENTRY
41.52

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL
ENTRY SESSION
-7.06 -7.06

TOTAL

SESSION 41.67

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# WEST

## **Generate Collection**

# Search Results - Record(s) 1 through 10 of 11 returned.

1. Document ID: US 20010006630 A1

L1: Entry 1 of 11

File: PGPB

Jul 5, 2001

PGPUB-DOCUMENT-NUMBER: 20010006630 PGPUB-FILING-TYPE: new-utility

DOCUMENT-IDENTIFIER: US 20010006630 A1

TITLE: INTRODUCING A BIOLOGICAL MATERIAL INTO A PATIENT

PUBLICATION-DATE: July 5, 2001

INVENTOR-INFORMATION:

NAME

CITY

STATE

RULE-47

YACOBY-ZEEVI, ORON

MEITAR

тт

COUNTRY

US-CL-CURRENT: 424/93.2; 424/94.64, 424/94.67, 435/325

| Full | Title | Citation | Front | Review | Classification | Date | Reference | Claims | KWC | Draw, Desc | Image |
|------|-------|----------|-------|--------|----------------|------|-----------|--------|-----|------------|-------|
|      | _     |          |       |        |                |      |           |        |     |            |       |

## 2. Document ID: US 6242238 B1

L1: Entry 2 of 11

File: USPT

Jun 5, 2001

US-PAT-NO: 6242238

DOCUMENT-IDENTIFIER: US 6242238 B1

TITLE: Isolated nucleic acid molecule encoding mammalian endoglucuronidase and

uses therefor

DATE-ISSUED: June 5, 2001

INVENTOR-INFORMATION:

STATE ZIP CODE COUNTRY CITY NAME AUX N/A N/A Freeman; Craig Geoffrey Rivett N/A AUX Hulett; Mark Darren Cook N/A Parish; Christopher Richard Campbell N/A N/A AUX Hamdorf; Brenton James Swinger Hill N/A N/A AUX

US-CL-CURRENT: 435/200; 435/252.3, 435/320.1, 536/23.1, 536/23.2

Full Title Citation Front Review Classification Date Reference Claims KMC Draw. Desc Image

☐ 3. Document ID: US 6190875 B1

L1: Entry 3 of 11 File: USPT Feb 20, 2001

US-PAT-NO: 6190875

DOCUMENT-IDENTIFIER: US 6190875 B1

TITLE: Method of screening for potential anti-metastatic and anti-inflammatory

agents using mammalian heparanase as a probe

DATE-ISSUED: February 20, 2001

INVENTOR-INFORMATION:

| NAME                   | CITY     | STATE | ZIP CODE | COUNTRY |
|------------------------|----------|-------|----------|---------|
| Ben-Artzi; Hanna       | Zion     | N/A   | N/A      | ILX     |
| Ayal-Hershkovitz; Maty | Herzliya | N/A   | N/A      | ILX     |
| Vlodavsky; Israel      | Zion     | N/A   | N/A      | ILX     |
| Pecker; Iris           | Zion     | N/A   | N/A      | ·ILX    |
| Peleg; Yoav            | Rehovot  | N/A   | N/A      | ILX     |
| Miron; Daphna          | Rehovot  | N/A   | N/A      | ILX     |

US-CL-CURRENT: 435/18; 435/201



KWC | Draw Desc | Image |

# 4. Document ID: US 6177545 B1

L1: Entry 4 of 11 File: USPT

PT Jan 23, 2001

US-PAT-NO: 6177545

DOCUMENT-IDENTIFIER: US 6177545 B1

TITLE: Heparanase specific molecular probes and their use in research and

medical applications

DATE-ISSUED: January 23, 2001

INVENTOR-INFORMATION:

| NAME              | CITY           | STATE | ZIP CODE | COUNTRY |
|-------------------|----------------|-------|----------|---------|
| Pecker; Iris      | Rishon Le Zion | N/A   | N/A      | ILX     |
| Vlodavsky; Israel | Mevaseret Zion | N/A   | N/A      | ILX     |
| Friedman; Yael    | Jerusalem      | N/A   | N/A      | ILX     |
| Perets; Tuvia     | Ramat Gan      | N/A   | N/A      | ILX     |

US-CL-CURRENT: 530/387.3; 530/388.1, 530/388.2, 530/388.26, 530/388.85, 530/389.1, 530/413

| Full Title | Citation | Front | Review | Classification | Date | Reference |
|------------|----------|-------|--------|----------------|------|-----------|

KWIC Draw Desc Image

# ☐ 5. Document ID: US 5968822 A

L1: Entry 5 of 11

File: USPT

Oct 19, 1999

US-PAT-NO: 5968822

DOCUMENT-IDENTIFIER: US 5968822 A

TITLE: Polynucleotide encoding a polypeptide having heparanase activity and

expression of same in transduced cells

DATE-ISSUED: October 19, 1999

INVENTOR-INFORMATION:

STATE ZIP CODE COUNTRY NAME CITY Pecker; Iris Rishon le Zion 75203 N/A N/A ILX Vlodavsky; Israel Mevaseret Zion 90805 N/A N/A ILX ILX Feinstein; Elena Rehovot 76214 N/A N/A

US-CL-CURRENT: 435/325; 435/200, 435/252.3, 435/320.1, 536/23.1, 536/23.2

Full Title Citation Front Review Classification Date Reference

KWIC Draw, Desc Image

# ☐ 6. Document ID: US 5362641 A

L1: Entry 6 of 11 File: USPT Nov 8, 1994

US-PAT-NO: 5362641

DOCUMENT-IDENTIFIER: US 5362641 A

TITLE: Heparanase derived from human Sk-Hep-1 cell line

DATE-ISSUED: November 8, 1994

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY Fuks; Zvi New York NY N/A N/A Vlodavsky; Israel Gilo N/A N/A ILX

US-CL-CURRENT: 435/209; 435/195, 435/200, 435/201

Full Title Citation Front Review Classification Date Reference KMC Draw Desc Image

7. Document ID: AU 200028786 A, WO 200052178 A1

L1: Entry 7 of 11 File: DWPI Sep 21, 2000

DERWENT-ACC-NO: 2000-579289

DERWENT-WEEK: 200065

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New polynucleotides encoding a polypeptide having heparanase activity, useful in wound healing and in gene therapy, particularly in treating tumor,

inflammation, autoimmunity, neurodegenerative diseases

INVENTOR: FEINSTEIN, E; PECKER, I; VLODAVSKY, I

PRIORITY-DATA: 1999US-0258892 (March 1, 1999)

PATENT-FAMILY:

 PUB-NO
 PUB-DATE
 LANGUAGE
 PAGES
 MAIN-IPC

 AU 200028786 A
 September 21, 2000
 N/A
 000
 C12N015/56

 WO 200052178 A1
 September 8, 2000
 E
 152
 C12N015/56

INT-CL (IPC): C12N 1/21; C12N 9/24; C12N 15/11; C12N 15/56; C12N 15/63

Full Title Citation Front Review Classification Date Reference

KWIC Draw Desc Image

# 8. Document ID: NO 200102190 A, WO 200025817 A1, AU 200013314 A

L1: Entry 8 of 11

File: DWPI

Jun 12, 2001

DERWENT-ACC-NO: 2000-399323

DERWENT-WEEK: 200141

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New monoclonal antibody comprising heparanase neutralizing activity, useful for treating a condition associated with heparanase expression e.g.

preventing angiogenesis

INVENTOR: AYAL-HERSHKOVITZ, M; FRIEDMAN, Y; MIRON, D; PECKER, I; PERETZ, T;

SHLOMI, Y ; VLODAVSKY, I

PRIORITY-DATA: 1998US-0186200 (November 4, 1998)

PATENT-FAMILY:

PUB-DATE LANGUAGE PAGES MAIN-IPC PUB-NO 000 A61K000/00 NO 200102190 A June 12, 2001 N/A May 11, 2000 Ε 028 A61K039/395 WO 200025817 A1 May 22, 2000 000 A61K039/395 AU 200013314 A N/A

INT-CL (IPC): A61K 0/00; A61K 38/47; A61K 39/395; C07K 16/40

Full Title Citation Front Review Classification Date Reference

KMC Draw Desc Image

## 9. Document ID: EP 1054980 A1, WO 9940207 A1, AU 9928319 A

L1: Entry 9 of 11

File: DWPI

Nov 29, 2000

DERWENT-ACC-NO: 1999-494300

DERWENT-WEEK: 200063

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New heparanase polypeptide useful for treating autoimmune diseases, skin

diseases, cardiovascular diseases and nervous system diseases including

Alzheimer's disease

INVENTOR: FUNAKUBO, M; NAKAJIMA, M; TOYOSHIMA, M

PRIORITY-DATA: 1998GB-0002725 (February 9, 1998)

PATENT-FAMILY:

| PUB-NO        | PUB-DATE          | LANGUAGE | PAGES | MAIN-IPC   |
|---------------|-------------------|----------|-------|------------|
| EP 1054980 A1 | November 29, 2000 | E        | 000   | C12N015/56 |
| WO 9940207 A1 | August 12, 1999   | E        | 040   | C12N015/56 |
| AU 9928319 A  | August 23, 1999   | N/A      | 000   | C12N015/56 |

INT-CL (IPC): A61K 31/70; A61K 38/47; A61K 39/395; C07K 16/40; C12N 5/10; C12N 9/24; C12N 15/56; G01N 33/50; G01N 33/573; G01N 33/577

| Full | Title | Citation | Front | Review | Classification | Date | Reference |
|------|-------|----------|-------|--------|----------------|------|-----------|

KWIC Draw. Desc Image

# 10. Document ID: AU 735116 B, WO 9911798 A1, AU 9891258 A, US 5968822 A, NO 9906228 A, EP 998569 A1, CZ 200000754 A3, HU 200002675 A2, CN 1272886 A

L1: Entry 10 of 11

File: DWPI

Jun 28, 2001

DERWENT-ACC-NO: 1999-302255

DERWENT-WEEK: 200142

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New human polynucleotide useful for treating angiogenesis, restenosis, and inflammation

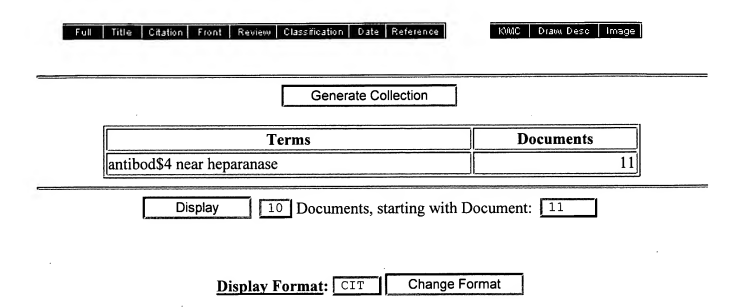
INVENTOR: FEINSTEIN, E; PECKER, I; VLODAVSKY, I

PRIORITY-DATA: 1998US-0109386 (July 2, 1998), 1997US-0922170 (September 2, 1997)

PATENT-FAMILY:

| PUB-NO          | PUB-DATE          | LANGUAGE | PAGES | MAIN-IPC   |
|-----------------|-------------------|----------|-------|------------|
| AU 735116 B     | June 28, 2001     | N/A      | 000   | C12N015/56 |
| WO 9911798 A1   | March 11, 1999    | E        | 054   | C12N015/56 |
| AU 9891258 A    | March 22, 1999    | N/A      | 000   | N/A        |
| US 5968822 A    | October 19, 1999  | N/A      | 000   | C12N015/56 |
| NO 9906228 A    | February 28, 2000 | N/A      | 000   | C12N000/00 |
| EP 998569 A1    | May 10, 2000      | E        | 000   | C12N015/56 |
| CZ 200000754 A3 | August 16, 2000   | N/A      | 000   | C12N015/56 |
| HU 200002675 A2 | December 28, 2000 | N/A      | 000   | C12N015/56 |
| CN 1272886 A    | November 8, 2000  | N/A      | 000   | C12N015/56 |

INT-CL (IPC): A61K 38/47; C12N 0/00; C12N 1/21; C12N 5/10; C12N 9/24; C12N 15/11; C12N 15/56; C12N 15/63



## Generate Collection

# **Search Results -** Record(s) 11 through 11 of 11 returned.

# 11. Document ID: JP 3188691 B2, WO 9102977 A, AU 9063364 A, EP 487627 A1, JP 05504047 W, US 5362641 A, AU 654804 B, EP 487627 A4, EP 487627 B1, DE 69032406 E

L1: Entry 11 of 11

File: DWPI

Jul 16, 2001

DERWENT-ACC-NO: 1991-087405

DERWENT-WEEK: 200142

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TITLE: Purificn. of heparanaseon pressure-sensitive adhesive polymer contg. by cation exchange resin chromatography and affinity absorbent purificn. of

heparinase-contg. cell extract

INVENTOR: FUKS, Z; VLODAVSKY, I

PRIORITY-DATA: 1989US-0397554 (August 23, 1989), 1992US-0768900 (January 8,

1992)

#### PATENT-FAMILY:

| PUB-NO        | PUB-DATE          | LANGUAGE | PAGES | MAIN-IPC   |
|---------------|-------------------|----------|-------|------------|
| JP 3188691 B2 | July 16, 2001     | N/A      | 020   | C12N009/24 |
| WO 9102977 A  | March 7, 1991     | N/A      | 000   | N/A        |
| AU 9063364 A  | April 3, 1991     | N/A      | 000   | N/A        |
| EP 487627 A1  | June 3, 1992      | E        | 000   | G01N033/53 |
| JP 05504047 W | July 1, 1993      | N/A      | 000   | C12N009/24 |
| US 5362641 A  | November 8, 1994  | N/A      | 023   | C12N009/26 |
| AU 654804 B   | November 24, 1994 | N/A      | 000   | C12N009/88 |
| EP 487627 A4  | March 3, 1993     | N/A      | 000   | N/A        |
| EP 487627 B1  | June 10, 1998     | E        | 000   | C12N009/24 |
| DE 69032406 E | July 16, 1998     | N/A      | 000   | C12N009/24 |

INT-CL (IPC): A61K 37/48; A61K 37/54; A61K 37/56; A61K 38/46; A61P 17/02; C07K 1/14; C12N 9/14; C12N 9/24; C12N 9/26; C12N 9/42; C12N 9/88; C12Q 1/34; G01N 33/48; G01N 33/53; G01N 33/573

Full Title Citation Front Review Classification Date Reference

KMC Draw Desc Image

## Generate Collection

| Terms                      | Documents |
|----------------------------|-----------|
| antibod\$4 near heparanase | 11        |

Display

10 Documents, starting with Document: 11

Display Format: CIT Change Format

# WEST

## **Generate Collection**

## **Search Results -** Record(s) 1 through 10 of 11 returned.

# ☐ 1. Document ID: US 20010006630 A1

L1: Entry 1 of 11

File: PGPB

Jul 5, 2001

DOCUMENT-IDENTIFIER: US 20010006630 A1

TITLE: INTRODUCING A BIOLOGICAL MATERIAL INTO A PATIENT

#### DRTX:

[0055] FIGS. 2a-b demonstrate that heparanase adheres to BMSCs and retains its activity. Cells that were incubated with heparanase were washed, collected and subjected to the (2a) DMB heparanase activity assay (1-6 represent six different experiments) and (2b) Western blot analysis using anti\_heparanase antibodies. T=Trypsin, 1E=1 mM EDTA, 2E=2 mM EDTA, Cb=control, purified heparanase from baculovirus, p60, Cc= control, purified heparanase from CHO cells, p45, kDa=kiloDaltons.

#### DRTX:

[0057] FIGS. 4a-c demonstrate that heparanase adheres to B16-F1 cells and retain its activity. Cells that were either transfected with the hpa cDNA ("transfected"), or incubated with heparanase ("adhered", +b22, or +b27), or not treated with heparanase (NT or -), were washed, collected and subjected to the DMB heparanase activity assay (4a), gel shift assay (4b), and Western blot analysis using anti heparanase antibodies (4c). Purified baculovirus heparanase p60 (b22, b27), or CHO heparanase p45 were used as controls (C).

## DRTX:

[0058] FIGS. 5a-b demonstrate that heparanase binds to CHO-dhfr cell line, undergoes proteolytic cleavage and exhibits high heparanase activity. Cells that were incubated with heparanase were washed, collected and subjected to DMB activity assay (5a), and Western blot analysis using anti-heparanase antibodies (5b).

## DETX:

[0126] Sputum viscosity and proteolytic activation of heparanase by sputum-borne proteases: 250 .mu.l of sputum samples, kept at 37.degree. C., were mixed in eppendorf tubes with either recombinant heparanase (p60), or with saline, or with a cocktail of protease inhibitors followed by the addition of heparanase, to make a total volume of 350 .mu.l. The samples were immediately transferred to 0.5 insulin syringes and tested for viscosity using a microviscosometer (Haake). The samples in the syringes were then incubated at 37.degree. C. and tested again for viscosity after 10, 50 and 120 minutes. Then, the samples were centrifuged for 10 minutes at 13,000 rpm and the supernatants were subjected to Western blot analysis, using several anti-heparanase antibodies (monoclonal Nos. 117 and 239, described in U.S. patent application Ser. No. 09/071,739, filed May 1, 1998).

Full Title Citation Front Review Classification Date Reference

KMC Draw Desc Image

# 2. Document ID: US 6242238 B1

L1: Entry 2 of 11

File: USPT

Jun 5, 2001

DOCUMENT-IDENTIFIER: US 6242238 B1

 ${\tt TITLE:}$  Isolated nucleic acid molecule encoding mammalian endoglucuronidase and uses therefor

#### DEPR:

An ELISA assay was developed for assaying for anti-human heparanase antibodies. The assay involved immobilising human platelet heparanase (5 .mu.g/ml in PBS, 15 hr, 4.degree. C.), purified from human platelets as previously described, in 96 well plastic microplates (25 .mu.l/well). Non-specific binding sites were then blocked by the addition of 200 .mu.l/well of PBS containing 1% (w/v) bovine serum albumin (BSA) for 2 hr at 4.degree. C. Following three washes with 200 .mu.l/well of PBS/0/05% Tween 20 (PBST), 50 .mu.l/well of serial dilutions of the antisera in PBS/1% BSA were added and incubated for 2 hr at 4.degree. C. Following three washes with PBST, 50 .mu.L/well of horse radish peroxidase (HRP) coupled sheep anti-rabbit Ig was added in PBS/1% BSA for 1 hr at 4.degree. C., the plate again washed three times with PBST, and bound HRP measured by the addition of the colourometric HRP substrate 2,2'-azino-bis (3-ethylbenthiazoline-6-sulfonic acid diammonium salt (ABTS), colour development being measured at 405 nm on an ELISA plate reader after 30 minutes incubation at 37.degree. C.



KWMC Draw. Desc Image

# 3. Document ID: US 6190875 B1

L1: Entry 3 of 11

File: USPT

Feb 20, 2001



DOCUMENT-IDENTIFIER: US 6190875 B1

TITLE: Method of screening for potential anti-metastatic and anti-inflammatory agents using mammalian heparanase as a probe

#### BSPR:

According to still further features in the described preferred embodiments the agent or agents include an anti-heparanase antibody.

#### DRPR

One example include anti-heparanase antibodies. It is well known that by binding to the active site antibodies can be used to inhibit catalytic activity of an enzyme.

#### DRPR:

Anti-heparanase antibodies are described in length in U.S. patent application Ser. No. 09/071,739, which is incorporated by reference as if fully set forth herein.

#### CLPR:

6. The method of claim 1, wherein the agent is an anti-heparanase antibody.

#### CLPR:

18. The method of claim 14, wherein the agent is an anti-heparanase antibody.

#### CLPR:

38. The quantitative method of claim 26, wherein the agent is an anti-heparanase antibody.

#### CLPR:

56. The quantitative method of claim 44, wherein the agent is an anti-heparanase antibody.

Full Title Citation Front Review Classification Date Reference

KMC Draw. Desc Image

## 4. Document ID: US 6177545 B1

L1: Entry 4 of 11

File: USPT

Jan 23, 2001

DOCUMENT-IDENTIFIER: US 6177545 B1

TITLE: Heparanase specific molecular probes and their use in research and medical applications

## BSPR:

The present invention relates to heparanase specific molecular probes their use in research and medical applications. More particularly, the present invention relates to the use of heparanase specific molecular probes, such as anti-heparanase antibodies (both poly- and monoclonal) and heparanase gene (hpa) derived nucleic acids, including, but not limited to, PCR primers, antisense oligonucleotide probes, antisense RNA probes, DNA probes and the like for detection and monitoring of malignancies, metastasis and other non-malignant conditions, efficiency of therapeutic treatments, targeted drug delivery and therapy.

## BSPR:

Heparanase activity could not be detected in normal stromal fibroblasts, mesothelial, endothelial and smooth muscle cells derived from non cancerous biopsies and effusions (12). These observations indicate that heparanase

expression may serve as a marker for tumor cells and in particular for those which are highly invasive or potentially invasive. If the same conclusion can be reached by immunostaining of tissue specimens, anti-heparanase antibodies may be applied for early detection and diagnosis of metastatic cell populations and micro-metastases.

#### RSPR

Expression of heparanase by cells of the immune system: Heparanase activity correlates with the ability of activated cells of the immune system to leave the circulation and elicit both inflammatory and autoimmune responses. Interaction of platelets, granulocytes, T and B lymphocytes, macrophages and mast cells with the subendothelial ECM is associated with degradation of heparan sulfate (HS) by heparanase activity (7). The enzyme is released from intracellular compartments (e.g., lysosomes, specific granules) in response to various activation signals (e.g., thrombin, calcium ionophore, immune complexes, antigens, mitogens), suggesting its regulated involvement and presence in inflammatory sites and autoimmune lesions. Heparan sulfate degrading enzymes released by platelets and macrophages are likely to be present in atherosclerotic lesions (21). Hence, cDNA probes and anti-heparanase antibodies may be applied for detection and early diagnosis of these lesions.

#### BSPR

On the basis of the examples described below, it appears that cDNA and RNA probes, PCR primers, and anti-heparanase antibodies (heparanase specific molecular probes) can be applied to detect the heparanase gene and protein and hence for early diagnosis of micrometastases, autoimmune lesions, renal failure and atherosclerotic lesions using biopsy specimens, plasma samples, and body fluids.

#### BSPR:

Collectively, it is evident that so far no one had succeeded in eliciting anti-heparanase antibodies.

#### BSPR

According to still further features in the described preferred embodiments the elicitation is through in vivo or in vitro techniques, the antibody having been prepared by a process comprising the steps of (a) exposing cells capable of producing antibodies to the heparanase protein or the immonogenical part thereof and thereby generating antibody producing cells; (b) fusing the antibody producing cells with myeloma cells and thereby generating a plurality of hybridoma cells each producing monoclonal antibodies; and (c) screening the plurality of monoclonal antibodies to identify a monoclonal antibody which specifically binds heparanase.

#### BSPR:

According to still further features in the described preferred embodiments the detectable heparanase specific molecular probe is selected from the group consisting of a nucleic acid sequence hybridizable with heparanase encoding nucleic acid and an anti-heparanase antibody capable of specifically binding heparanase.

#### BSPR:

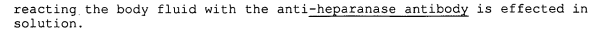
According to further features in preferred embodiments of the invention described below, there is provided a method of detecting heparanase protein in a body fluid of a patient comprising the steps of reacting the body fluid with an anti-heparanase antibody and monitoring the reaction.

## BSPR:

According to still further features in the described preferred embodiments the anti-heparanase antibody is selected from the group consisting of a monoclonal antibody and a poly clonal antibody.

#### BSPR:

According to still further features in the described preferred embodiments



#### BSPR:

According to still further features in the described preferred embodiments reacting the body fluid with the anti-heparanase antibody is effected on a substrate capable of adsorbing proteins present in the body fluid.

#### BSPR:

According to further features in preferred embodiments of the invention described below, there is provided a method of detecting the presence, absence or level of heparanase protein in a biological sample comprising the steps of (a) extracting proteins from the biological sample, thereby obtaining a plurality of proteins; (b) size separating the proteins; (c) interacting the size separated proteins with an anti-heparanase antibody; and (d) detecting the presence, absence or level of the interacted anti-heparanase antibody.

#### BSPR:

According to still further features in the described preferred embodiments the anti-heparanase antibody is selected from the group consisting of a polyclonal antibody and a monoclonal antibody.

#### BSPR:

According to further features in preferred embodiments of the invention described below, there is provided a method of targeted drug delivery to a tissue of a patient, the tissue expressing heparanase, the method comprising the steps of providing a complex of a drug directly or indirectly linked to an anti-heparanase antibody and administering the complex to the patient.

#### BSPR:

According to further features in preferred embodiments of the invention described below, there is provided a method of treating a patient having a condition associated with heparanase expression comprising the step of administering an anti-heparanase antibody to the patient.

## DRPR:

FIGS. 17a-b demonstrate Western blots of extracts of cells expressing various segments of heparanase as detected with polyclonal anti heparanase antibodies. 17a--antiserum from rabbit 7640, 17b--antiserum from rabbit 7644. Lane 1, E. coli BL21(DE3)pLysS cells transfected with pRSET, lane 2, E. coli BL21(DE3)pLysS cells transfected with pRSET containing the heparanase entire open reading frame (543 amino acids, SE ID NOs: 2 and 3), lane 3, E. coli BL21(DE3)pLysS cells transfected with pRSEThpaBK containing 414 amino acids of the heparanase open reading frame (amino acids 130-543 of SEQ ID NOS: 2 and lane 4, E. coli BL21(DE3)pLysS cells transfected with pRSEThpaBH containing 302 amino acids of the heparanase open reading frame (amino acids 130-431 of SEQ ID NOs: 2 and 3), lane 5, molecular size markers, lane 6, medium of Sf21 insect cells infected with recombinant Baculovirus pFhpa containing the heparanase entire open reading frame (543 amino acids, SEQ ID NOs: 2 and 3), lane 7, Sf21 insect cells infected with recombinant baculovirus with no insert. Proteins were separated on 10% SDS-PAGE, antisera were diluted 1:1,000. Detection was performed by ECL (Amersham) according to the manufacturer's instructions. Size in kDa is shown to the right, as was determined using prestained SDS-PAGE standards, Bio-Rad, CA..

#### DRPR:

FIG. 18 demonstrates Western blot using affinity purified polyclonal antibodies with heparanase expressed in various expression systems. Lane 1, medium of Sf21 insect cells infected with recombinant Baculovirus pFhpa, lane 2, cell extract of a Chinese hamster ovary (CHO) clone stably transfected with a vector containing no insert, lane 3, cell extract of a CHO stable clone transfected with hpa cDNA, lane 4, proteins precipitated from medium of the yeast Pichia pastoris transfected with hpa cDNA. Proteins were separated on 4-20% gradient SDS-PAGE, antibody was diluted 1:100. Detection was performed by ECL (Amersham) according to the manufacturer's instructions. For CHO and

Pichia clones see U.S. patent application Ser. No. 09/071,618, entitled "RECOMBINANT CELLS AND METHODS FOR EXPRESSING RECOMBINANT HEPARANASE AND METHOD OF PURIFYING SAME", filed on the date of filing of the present application, which is incorporated by reference as if fully set forth herein. Size in kDa is shown to the right, as was determined using prestained SDS-PAGE standards, Bio-Rad, CA..

#### DEPR:

The present invention is of heparanase specific molecular probes which can be used in research and medical applications. Specifically, the present invention can be used for the detection and monitoring of malignancies, metastasis and other, non-malignant conditions, efficiency of therapeutic treatments, targeted drug delivery and therapy, using heparanase specific molecular probes, such as anti-heparanase antibodies (both poly- and monoclonal) and heparanase gene (hpa) derived nucleic acids, including, but not limited to, PCR primers, antisense oligonucleotide probes, antisense RNA probes, DNA probes and the like.

#### DEPR:

As used herein in the specification and in the claims section below, the term "detectable heparanase specific molecular probe" and its equivalent term "detectable heparanase molecular probe" both refer to a nucleic acid sequences hybridizable with heparanase encoding nucleic acid or to an anti-heparanase antibody capable of specifically binding heparanase. The nucleic acid sequence hybridizable with heparanase encoding nucleic acid is, for example, a synthetic oligonucleotide, an antisesnse heparanase RNA or heparanase DNA, and it is preferably labeled by the detectable moiety.

#### DEPR:

Therefore, according to another aspect of the present invention there is provided a method of detecting heparanase protein in a body fluid of a patient. The method comprises the steps of reacting the body fluid with an anti-heparanase antibody, either poly or monoclonal antibody, and monitoring the reaction. The body fluid is, for example, plasma, urine, pleural effusions or saliva. Monitoring the reaction may be effected by having the antibody labeled with a detectable moiety, or to use its constant region as an inherent detectable moiety, to which a second antibody which includes a detectable moiety can specifically bind.

## DEPR:

According to a preferred embodiment of the present invention reacting the body fluid with the anti-heparanase antibody is effected in solution. Alternatively, reacting the body fluid with the anti-heparanase antibody is effected on a substrate capable of adsorbing proteins present in the body fluid, all as well known in the art of antibody based diagnosis.

#### DEPR:

As further shown in the Examples section below, protein blots and anti-heparanase antibodies prove useful in detecting the presence, absence or level of heparanase protein in various biological samples.

## DEPR:

Therefore, further according to the present invention there is provided a method of detecting the presence, absence or level of heparanase protein in a biological sample. The method comprises the following steps. First, proteins are extracted from the biological sample, thereby a plurality of proteins are obtained. The protein extract may be a crude extract and can also include non-proteinacious material. Second, the proteins are size separated, e.g., by electrophoresis, gel filtration etc. Fourth, the size separated proteins are interacted with an anti-heparanase antibody, either poly or monoclonal antibody. Finally, the presence, absence or level of the interacted anti-heparanase antibody is detected. In case of gel electrophoresis the interaction with the antibody is typically performed following blotting of the size separated proteins onto a solid support (membrane).

#### DEPR:

Therefore, according to yet another aspect of the present invention there is provided a method of targeted drug delivery to a tissue of a patient, the tissue expressing heparanase. The method comprises the steps of providing a complex of a drug directly or indirectly linked to an anti-heparanase antibody and administering the complex to the patient. External radio imaging is also envisaged, wherein the drug is replaced with an imageable radio isotope. Endoscopic or laparoscopic imaging is also envisaged. In the latter cases the drug is typically replaced by a fluorescence or luminescence substance. These procedures may, for example, be effective in finding/destroying micrometastases.

#### DEPR:

Therefore, according to another aspect of the present invention there is provided a method of treating a patient having a condition associated with heparanase expression. The method comprises the step of administering an anti-heparanase antibody to the patient.

#### DEPR:

Preferably, the elicitation of the antibody is through in vivo or in vitro techniques, the antibody having been prepared by a process comprising the steps of, first, exposing cells capable of producing antibodies to the <a href="heparanase">heparanase</a> protein or the immonogenical part thereof and thereby generating antibody producing cells. second, fusing the antibody producing cells with myeloma cells and thereby generating a plurality of hybridoma cells each producing monoclonal antibodies, and third, screening the plurality of monoclonal antibodies to identify a monoclonal antibody which specifically binds heparanase.

#### DEPR:

In situ detection of heparanase by antibodies: hpa-transfected and non transfected CHO cells were plated on 8-chamber tissue culture slides (Nunc). Cells were fixed in 95% ethanol, 5% acetic acid for 5 minutes at -20.degree. C. Cells were permeabilized using permeabilization buffer (20 mM HEPES, pH 7.4; 300 mM Sucrose; 50 mM NaCl; 3 mM MgCl.sub.2; 0.5% Triton X-100) for 4 minutes at 4.degree. C. Endogenous peroxidases were blocked using 0.3% H.sub.2 O.sub.2 in methanol and non specific binding sites were blocked using 5% horse serum in PBS. Monoclonal anti-heparanase antibody (supernatant of hybridoma) was applied and incubated with the cells overnight at room temperature. Antibody was washed away and biotinylated secondary antibody (horse-anti mouse, Vector, Vectastain ABC system) was added for 30 minutes at roomtemperature. Immunostaining was detected using Di Amino Benzidine and H.sub.2 O.sub.2 (Sigma tablets) until desired staining-intensity was achieved. Slides were counterstained with Mayer's hematoxylin. Immunostaining with polyclonal antibodies was performed under the same conditions, affinity purified antibody was used at 1:500 dilution. Biotinylated horse anti-rabbit was used as a secondary antibody (Vector, Vectastain ABC system). Blood smears were prepared from a healthy donor. Fixation and staining were performed as described above.

## DEPR:

This is the first result suggesting a role for heparanase in the pathogenesis of proteinuria in type I diabetes. Obviously, measurements of urinary heparanase activity is both time consuming and not sensitive enough. Moreover, we have demonstrated the presence of an inhibitor of mammalian heparanase in the urine of normal individuals. The nature of this inhibitory substance, possibly urinary glycosaminoglycans is currently being studied. Urinary heparanase activity is therefore the result of a balance between the presence in the urine of the enzyme and its inhibitor(s). Immunodetection of the heparanase protein is therefore a more sensitive and straightforward approach for diagnostic purposes. Altogether, our results clearly indicate that anti-heparanase antibodies that identify the heparanase antigen can be applied for early diagnosis of cancer metastasis and renal diseases. As discussed above, it is conceivable that heparanase may overcome the filtration barrier of the glomerular basement membrane and ECM simply by virtue of its ability to

degrade the HS moieties that are held responsible for their permeaselective properties. Urinary heparanase is therefore expected to reflect the presence of heparanase in the circulation and hence be a sensitive marker for metastatic, inflammatory and kidney disease. Of particular significance is the potential ability to follow the course of tumor progression and spread, response to anti-cancer treatments, and possible relapse of the disease in a given patient. Targeted drug delivery and therapy are another aspect of the use for such antibodies.

#### DEPR:

Availability of anti-heparanase antibodies will enable development of immunological assays for screening tissue and body fluids for heparanase. An ELISA will provide a more sensitive and convenient means of detection as compared to the currently available assays of heparanase activity which do not appear sensitive enough for the detection of the enzyme in non-concentrated plasma and body fluids.

| Full | Title | Citation | Front | Review | Classification | Date | Reference |
|------|-------|----------|-------|--------|----------------|------|-----------|

KMC Draw Desc Image

## 5. Document ID: US 5968822 A

L1: Entry 5 of 11

File: USPT

Oct 19, 1999

DOCUMENT-IDENTIFIER: US 5968822 A

TITLE: Polynucleotide encoding a polypeptide having heparanase activity and expression of same in transduced cells

## BSPR:

Other potential therapeutic applications: Apart from its involvement in tumor cell metastasis, inflammation and autoimmunity, mammalian heparanase may be applied to modulate: bioavailability of heparin-binding growth factors (5); cellular responses to heparin-binding growth factors (e.g., bFGF, VEGF) and cytokines (IL-8) (31a, 29); cell interaction with plasma lipoproteins (32); cellular susceptibility to certain viral and some bacterial and protozoa infections (33, 33a, 33b); and disintegration of amyloid plaques (34). Heparanase may thus prove useful for conditions such as wound healing, angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases and viral infections. Mammalian heparanase can be used to neutralize plasma heparin. as a potential replacement of protamine. Anti-heparanase antibodies may be applied for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Common use in basic research is expected.

#### DEPR:

Anti-heparanase antibodies, which may be raised against the recombinant enzyme, would be useful for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Such antibodies may also serve as neutralizing agents for heparanase activity.

#### DEPR:

The recombinant protein may be purified by any conventional protein purification procedure close to homogeneity and/or be mixed with additives. The recombinant protein may be manufactured using any of the cells described above. The recombinant protein may be in any form. It may be in a crystallized form, a dehydrated powder form or in solution. The recombinant protein may be useful in obtaining pure heparanase, which in turn may be useful in eliciting anti-heparanase antibodies, either poly or monoclonal antibodies. and as a screening active ingredient in an anti-heparanase inhibitors or drugs screening assay or system.

Full Title Citation Front Review Classification Date Reference

KWC Draw, Desc Image

☐ 6. Document ID: US 5362641 A

L1: Entry 6 of 11

File: USPT

Nov 8, 1994

DOCUMENT-IDENTIFIER: US 5362641 A

TITLE: Heparanase derived from human Sk-Hep-1 cell line

#### DRPR:

FIG. 6: Preparative native PAGE. Active fractions eluted from Con-A-Sepharose were pooled, dialyzed and applied to a native 8% polyacrylamide gel. The gel was cut into 5 mm strips (0-8) and heparanase activity in material electroeluted from each strip was assayed by gel filtration on Sepharose 6B of labeled degradation products released from sulfate labeled ECM. Material associated with strip #7 was injected into rabbits to produce the polyclonal anti-heparanase antibodies used in FIGS. 5AI through 5DII:

#### DRPR:

FIGS. 7A and 7B: Identification of heparanase in extracts of a biopsy specimen from a human ovarian tumor. Ovarian tumor removed at surgery was homogenized and the supernatant fraction subjected to FPLC gel filtration on superose 12 column. The starting material (lane 1) and fractions number 16-19 of the active peak (lanes 2-5) were subjected to SDS/PAGE and "Western" blot analysis. FIG. 7A: Coomassie blue staining of proteins electrotransferred to Immobilon-P transfer membrane. FIG. 7B: Autoradiogram of the same transfer membrane following successive incubations with rabbit anti-heparanase antibodies and .sup.125 I-goat anti-rabbit IgG.

### DEPR:

In order to obtain a single band preparation, enzyme eluted from Con A-Sepharose was subjected to native polyacrylamide gel electrophoresis, as described in Materials and Methods. The gel was cut into strips of 5 mm, and protein was electroeluted from 5 mm segments of each strip. Heparanase activity, measured by gel filtration analysis of sulfate-labelled degradation products, was eluted primarily from strip #7. A much lower activity was detected in strip #4 (FIG. 6). Enzyme associated with strip #7 was homogenized with the polyacrylamide in a minimal volume of PBS, mixed with complete Freund's adjuvant, and injected into rabbits to produce polyclonal anti-heparanase antibodies. This material will also be subjected to amino acid sequencing for the purpose of gene cloning and expression. The anti-heparanase antibodies have been used to immunodetect the enzyme in "Western" blots of fractions eluted from CD-Sephadex (FIG. 5AII(B)) and Con-A Sepharose and in active fractions derived from a biopsy specimen of a human ovarian tumor (FIGS. 7A and 7B).

Full Title Citation Front Review Classification Date Reference

KWMC Draw Desc Image

7. Document ID: AU 200028786 A, WO 200052178 A1

L1: Entry 7 of 11

File: DWPI

Sep 21, 2000

DERWENT-ACC-NO: 2000-579289

DERWENT-WEEK: 200065

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New polynucleotides encoding a polypeptide having heparanase activity, useful in wound healing and in gene therapy, particularly in treating tumor, inflammation, autoimmunity, neurodegenerative diseases

## ABTX:

in vivo eliciting of anti-heparanase antibodies comprising administering the nucleic acid construct of (1) including a segment of the novel polynucleotide; and

#### ABTX:

(11) a DNA vaccine for eliciting in vivo anti-heparanase antibodies comprising the nucleic acid construct of (1) and a promoter for directing the expression of the polynucleotide segment in vivo.

#### ABTX:

USE - The polynucleotide is useful in gene therapy, particularly in treating tumor, inflammation or autoimmunity. Particularly, the polynucleotide is useful in modulating the bioavailability of heparin-binding growth factors, cellular responses to heparin-binding growth factors (e.g. bFGF) and cytokines (e.g. interleukin (IL)-8), cell interaction with plasma lipoproteins, cellular susceptibility to certain viral and some bacterial and protozoa infections, or disintegration of neurodegenerative plaques. The polynucleotide is useful in wound healing (e.g. thermal, chemical or radiation burns), and in the treatment of angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases (Gerstmann-Straussler Syndrome or Creutzfeldt-Jakob disease), and some viral, bacterial or protozoa infections. The polynucleotide is also useful in developing diagnostic assays for these diseases, for providing new tools for basic research especially in the field of medicine and biology, and for developing new drugs to inhibit these diseases. The nucleic acid constructs or vectors, antisense oligonucleotides, and ribozymes are useful for modulating heparanase. The polynucleotide sequences can be used to provide DNA vaccines that will elicit in vivo anti-heparanase antibodies. The polypeptide is useful for catalyzing the degradation of heparan sulfate in an in vitro assay. The recombinant proteins are also useful for obtaining pure heparanase, which in turn may be useful in eliciting anti-heparanase antibodies, either poly- or monoclonal antibodies (for immunodetection), and as a screening active ingredient in an anti-heparanase inhibitors or drugs screening assay or system.

Full Title Citation Front Review Classification Date Reference

KWC Drawl Desc Image

8. Document ID: NO 200102190 A, WO 200025817 A1, AU 200013314 A

L1: Entry 8 of 11

File: DWPI

Jun 12, 2001

DERWENT-ACC-NO: 2000-399323

DERWENT-WEEK: 200141

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New monoclonal antibody comprising heparanase neutralizing activity, useful for treating a condition associated with heparanase expression e.g. preventing angiogenesis

## ABTX:

NOVELTY - Treating a condition associated with heparanase expression comprises administering an anti-heparanase monoclonal <u>antibody having heparanase</u> neutralizing catalytic activity.

Full Title Citation Front Review Classification Date Reference

KWC Draw Desc Image

# 9. Document ID: EP 1054980 A1, WO 9940207 A1, AU 9928319 A

L1: Entry 9 of 11

File: DWPI

Nov 29, 2000

DERWENT-ACC-NO: 1999-494300

DERWENT-WEEK: 200063

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New heparanase polypeptide useful for treating autoimmune diseases, skin diseases, cardiovascular diseases and nervous system diseases including Alzheimer's disease

#### ABTX:

USE - The heparanase protein or the anti-heparanase antibody are used in pharmaceutical compositions for treating warm blooded animals suffering from a disease resulting from shortage/lack of the heparanase protein, or from excessive activity/over-expression of the heparanase protein, respectively. The heparanase protein is used in treating diseases such as trauma, autoimmune disease, skin diseases, cardiovascular diseases and nervous system diseases including Alzheimer's disease resulting from shortage or lack of polypeptide.

#### ABTX:

The anti-heparanase antibody is used in treating the diseases like cancer, cancer metastasis, angiogenesis and inflammation including arthritis resulting from excessive activity or over expression of heparanase protein. The polynucleotide coding for the polypeptide is used in gene therapy strategy (claimed). The anti-heparanase antibody can be used to detect the presence or absence of polypeptide and its concentration. Hence the anti-heparanase antibodies are used as diagnostic markers for the disorders such as cancer, cancer metastasis and angiogenesis. Agonist and antagonist can replace heparanase protein and anti-heparanase antibody respectively to treat the diseases caused by increased or decreased expression of heparanase protein.

Full Title Citation Front Review Classification Date Reference

KMC Draw Desc Image

10. Document ID: AU 735116 B, WO 9911798 A1, AU 9891258 A, US 5968822 A, NO 9906228 A, EP 998569 A1, CZ 200000754 A3, HU 200002675 A2, CN 1272886 A

L1: Entry 10 of 11

File: DWPI

Jun 28, 2001

DERWENT-ACC-NO: 1999-302255

DERWENT-WEEK: 200142

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New human polynucleotide useful for treating angiogenesis, restenosis, and inflammation

#### ABTX:

USE - The recombinant protein is used as a modulator of heparin-binding growth factors, cellular responses to heparin-binding growth factors and cytokines, cell interaction with plasma lipoproteins, cellular susceptibility to viral, protozoal and bacterial infections or disintegration of neurodegenerative plaques (claimed). Heparanase may be useful for conditions such as wound healing, angiogenesis, restenosis, athersclerosis, inflammation, neurodegenerative diseases, and viral infections. Mammalian heparanase can be used to neutralize plasma heparin, and anti-heparanase antibodies may be applied for immunodetection and diagnosis of micrometastases, autoimmune lesions, and renal failure in biopsy specimens, plasma samples, and body fluids.

## ABEQ:

USE - The recombinant protein is used as a modulator of heparin-binding growth factors, cellular responses to heparin-binding growth factors and cytokines, cell interaction with plasma lipoproteins, cellular susceptibility to viral, protozoal and bacterial infections or disintegration of neurodegenerative plaques (claimed). Heparanase may be useful for conditions such as wound healing, angiogenesis, restenosis, athersclerosis, inflammation, neurodegenerative diseases, and viral infections. Mammalian heparanase can be used to neutralize plasma heparin, and anti-heparanase antibodies may be applied for immunodetection and diagnosis of micrometastases, autoimmune lesions, and renal failure in biopsy specimens, plasma samples, and body fluids.

| Full   Title   | Citation Front Review Classification Date Referer | nce KWMC Draww Desc   Image |  |
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